L Number	Hits	Search Text	DB	Time stamp
-	1	DmGPCR	USPAT;	2002/07/24 09:38
			US-PGPU	3;
			EPO; JPO;	
			DERWEN	Γ
-	75	drosophila same receptor same coupled	USPAT;	2002/07/24 09:38
		,	US-PGPUE	3;
			EPO; JPO;	
			DERWEN	Γ
-	11	drosophila same receptor same coupled same bind	USPAT;	2002/07/24 09:41
ii.			US-PGPUI	3;
			EPO; JPO;	
			DERWEN*	Γ

# 09693746 Results

```
RESULT
AAU03346
     AAU03346 standard; Peptide; 9 AA.
ХХ
AC
     AAU03346;
ХХ
DT
     12-SEP-2001 (first entry)
XX
DE
     Fruit fly G protein coupled receptors, DmGPCR6aL/bL ligand #94.
XX
     Fruit fly; G protein coupled receptor; DmGPCR6aL/bL;
KW
КW
     human immunodeficiency virus; HIV; cancer; Parkinson's disease;
ЕW
     diabetes; obesity; atherosclerosis; thrombosis; stroke; renal
failure;
КW
     inflammation; rheumatoid arthritis; autoimmune disorder;
KW
     neurological disorder; schizophrenia; manic depression; dementia;
     severe mental retardation; dyskinesia; Huntington's disease;
ΗW
     Tourette's syndrome; ligand.
XX
OS
     Drosophila melanogaster.
XX
FH
                     Location/Qualifiers
     Key
FΤ
     Modified-site
                     /note= "Optionally, Tyr has an attached SO3H
FT
moiety"
     Modified-site
FT
                     /note= "C-terminus is amidated"
FT
XX
PN
     WO200131005-A2.
XX
PD
     03-MAY-2001.
ΧХ
     20-OCT-2000; 2000WO-US29002.
PF
XX
PR
     22-OCT-1999;
                    99US-0425676.
XX
PΑ
     (PHAA ) PHARMACIA & UPJOHN CO.
XX
     Lowery DE, Smith VG, Kubiak TA, Larsen MJ;
PΙ
XX
DR
     WPI; 2001-316333/33.
XX
PΤ
     New Drosophila melanogaster GPCR nucleic acids and polypeptide
useful
PT
     for inducing an immune response, for identifying homologs and for
PΤ
     treating e.g. diabetes, obesity and manic depression -
XX
PS
     Example 9; Page 101; 110pp; English.
XX
     The sequence is a fruit fly G protein coupled receptors,
DmGPCR6aL/bL,
     peptide ligand. The proteins are useful for inducing an immune
CC
response
```

```
against itself in a mammal. The nucleic acids are useful for
identifying
     an animal homolog of DmGPCR, by screening databases or libraries.
CC
The
CC
     compounds identified as binding partners or modulators of GPCR
binding
     are useful for treating diseases in animals, and for control
insects that
     are harmful or cause injury to plants or animals. Diseases treated
CC
     include infections (e.g. viral and human immunodeficiency virus,
HIV),
     cancer, pain, Parkinson's disease, hypotension, hypertension,
CC
diabetes,
     obesity, atherosclerosis, thrombosis, stroke, renal failure,
CC
     inflammation, rheumatoid arthritis, autoimmune disorders, and
CC
psychotic
     and neurological disorders (anxiety, schizophrenia, manic
depression,
     delirium, dementia, severe mental retardation, dyskinesias,
Huntington's
     disease or Tourette's syndrome). The nucleic acids can be used for
     genetic mapping, and producing the GPCRs. Anti-GPCR antibodies can
CC
be
     used in therapy, diagnostic assays and for modulating GPCR
CC
activity.
XX
SQ
     Sequence
                9 AA;
                          100.0%; Score 54; DB 22; Length 9;
  Query Match
                          100.0%; Pred. No. 6.4e+05;
  Best Local Similarity
             9; Conservative 0; Mismatches
  Matches
                                                 0; Indels
                                                                 0;
Gaps
        1 FDDYGHLRF 9
Qу
          1 fddyghlrf 9
Db
RESULT
AAU03347
    AAU03347 standard; Peptide; 9 AA.
ID
XX
AC
    AAU03347;
XX
     12-SEP-2001 (first entry)
DT
XX
     Fruit fly G protein coupled receptors, DmGPCR6aL/bL ligand #95.
DΕ
XX
KW
     Fruit fly; G protein coupled receptor; DmGPCR6aL/bL;
     human immunodeficiency virus; HIV; cancer; Parkinson's disease;
KW
KW
     diabetes; obesity; atherosclerosis; thrombosis; stroke; renal
failure;
     inflammation; rheumatoid arthritis; autoimmune disorder;
KW
    neurological disorder; schizophrenia; manic depression; dementia;
KW
     severe mental retardation; dyskinesia; Huntington's disease;
KW
KW
    Tourette's syndrome; ligand.
```

XX OS Drosophila melanogaster. XXFΗ Key Location/Qualifiers FΤ Modified-site /note= "C-terminus is amidated" FT XXW0200131005-A2. PNXX03-MAY-2001. PD XX20-OCT-2000; 2000WO-US29002. ΡF XXPR 22-OCT-1999; 99US-0425676. XX(PHAA ) PHARMACIA & UPJOHN CO. PΑ XXPΙ Lowery DE, Smith VG, Kubiak TA, Larsen MJ; XX WPI; 2001-316333/33. DR XX New Drosophila melanogaster GPCR nucleic acids and polypeptide PTuseful for inducing an immune response, for identifying homologs and for PΤ treating e.g. diabetes, obesity and manic depression -PTXX PS Example 9; Page 101; 110pp; English. XX The sequence is a fruit fly G protein coupled receptors, CCDmGPCR6aL/bL, CCpeptide ligand. The proteins are useful for inducing an immune response CC against itself in a mammal. The nucleic acids are useful for identifying CC an animal homolog of DmGPCR, by screening databases or libraries. The CC compounds identified as binding partners or modulators of GPCR binding are useful for treating diseases in animals, and for control CC insects that are harmful or cause injury to plants or animals. Diseases treated include infections (e.g. viral and human immunodeficiency virus, CCHIV), CCcancer, pain, Parkinson's disease, hypotension, hypertension, diabetes, obesity, atherosclerosis, thrombosis, stroke, renal failure, CC inflammation, rheumatoid arthritis, autoimmune disorders, and psychotic and neurological disorders (anxiety, schizophrenia, manic CCdepression, delirium, dementia, severe mental retardation, dyskinesias, Huntington's CC disease or Tourette's syndrome). The nucleic acids can be used for genetic mapping, and producing the GPCRs. Anti-GPCR antibodies can CC be used in therapy, diagnostic assays and for modulating GPCR activity.

```
XX
SQ
     Sequence 9 AA;
  Query Match
                          100.0%; Score 54; DB 22; Length 9;
  Best Local Similarity 100.0%; Pred. No. 6.4e+05;
          9; Conservative 0; Mismatches 0; Indels
  Matches
                                                                 0;
Gaps
QУ
       1 FDDYGHLRF 9
          1 fddyghlrf 9
RESULT
AAU03351
     AAU03351 standard; Peptide; 9 AA.
XX
AC
    AAU03351;
XX
    12-SEP-2001 (first entry)
DT
XX
     Fruit fly G protein coupled receptor ligand, drosulfakinin-1.
DE
XX
     Fruit fly; G protein coupled receptor; drosulfakinin-1;
KW
KW
     human immunodeficiency virus; HIV; cancer; Parkinson's disease;
     diabetes; obesity; atherosclerosis; thrombosis; stroke; renal
failure;
     inflammation; rheumatoid arthritis; autoimmune disorder;
KW
KW
     neurological disorder; schizophrenia; manic depression; dementia;
КW
     severe mental retardation; dyskinesia; Huntington's disease;
KW
     Tourette's syndrome; ligand.
XX
os
     Drosophila melanogaster.
XX
FΗ
                    Location/Qualifiers
     Key
FT
     Modified-site
FT
                     /note= "Tyr has an attached SO3H moiety"
FT
     Modified-site
FT
                     /note= "C-terminus is amidated"
XX
PN
     WO200131005-A2.
XX
PD
     03-MAY-2001.
XX
     20-OCT-2000; 2000WO-US29002.
ΡF
XX
     22-OCT-1999;
PR
                   99US-0425676.
XX
PΑ
     (PHAA ) PHARMACIA & UPJOHN CO.
XX
PΙ
     Lowery DE, Smith VG, Kubiak TA, Larsen MJ;
XX
DR
     WPI; 2001-316333/33.
XX
PΤ
     New Drosophila melanogaster GPCR nucleic acids and polypeptide
useful
```

```
for inducing an immune response, for identifying homologs and for
     treating e.g. diabetes, obesity and manic depression -
PΤ
XX
     Example 9; Page 98; 110pp; English.
PS
XX
     The sequence is a fruit fly G protein coupled receptor ligand,
CC
CC
     drosulfakinin-1. The proteins are useful for inducing an immune
response
     against itself in a mammal. The nucleic acids are useful for
identifying
     an animal homolog of DmGPCR, by screening databases or libraries.
CC
The
     compounds identified as binding partners or modulators of GPCR
CC
binding
     are useful for treating diseases in animals, and for control
CC
insects that
     are harmful or cause injury to plants or animals. Diseases treated
CC
     include infections (e.g. viral and human immunodeficiency virus,
HIV).
    cancer, pain, Parkinson's disease, hypotension, hypertension,
CC
diabetes,
     obesity, atherosclerosis, thrombosis, stroke, renal failure,
     inflammation, rheumatoid arthritis, autoimmune disorders, and
psychotic
CC
     and neurological disorders (anxiety, schizophrenia, manic
depression,
     delirium, dementia, severe mental retardation, dyskinesias,
Huntington's
     disease or Tourette's syndrome). The nucleic acids can be used for
CC
     genetic mapping, and producing the GPCRs. Anti-GPCR antibodies can
CC
be
     used in therapy, diagnostic assays and for modulating GPCR
CC
activity.
XX
     Sequence
                9 AA;
SQ
  Query Match
                          100.0%; Score 54; DB 22; Length 9;
                          100.0%; Pred. No. 6.4e+05;
  Best Local Similarity
  Matches
            9; Conservative 0; Mismatches 0; Indels
                                                                 0;
Gaps
        1 FDDYGHLRF 9
Qу
          Db
        1 fddyghlrf 9
RESULT
AAU03353
ID
     AAU03353 standard; Peptide; 14 AA.
XX
AC
    AAU03353;
XX
DT
     12-SEP-2001 (first entry)
XX
     Fruit fly G protein coupled receptor ligand, drosulfakinin-2.
DΕ
XX
```

```
KW
     Fruit fly; G protein coupled receptor; drosulfakinin-2;
KW
     human immunodeficiency virus; HIV; cancer; Parkinson's disease;
     diabetes; obesity; atherosclerosis; thrombosis; stroke; renal
HW
failure;
     inflammation; rheumatoid arthritis; autoimmune disorder;
     neurological disorder; schizophrenia; manic depression; dementia;
КW
     severe mental retardation; dyskinesia; Huntington's disease;
КW
KW
     Tourette's syndrome; ligand.
XX
     Drosophila melanogaster.
OS
XX
FH
                     Location/Qualifiers
FΤ
     Modified-site
FT
                     /note= "Tyr has an attached SO3H moiety"
FT
     Modified-site
                     14
                     /note= "C-terminus is amidated"
FT
XX
PN
     WO200131005-A2.
XX
PD
     03-MAY-2001.
XX
     20-OCT-2000; 2000WO-US29002.
ΡF
XX
PR
     22-OCT-1999;
                    99US-0425676.
XX
PΑ
     (PHAA ) PHARMACIA & UPJOHN CO.
XX
PΙ
     Lowery DE, Smith VG, Kubiak TA, Larsen MJ;
ХX
DR
     WPI; 2001-316333/33.
ХX
PT
     New Drosophila melanogaster GPCR nucleic acids and polypeptide
useful
PT
     for inducing an immune response, for identifying homologs and for
PT
     treating e.g. diabetes, obesity and manic depression -
XX
PS
     Disclosure; Page 4; 110pp; English.
XX
CC
     The sequence is a fruit fly G protein coupled receptor ligand,
CC
     drosulfakinin-2. The proteins are useful for inducing an immune
response
     against itself in a mammal. The nucleic acids are useful for
CC
identifying
CC
     an animal homolog of DmGPCR, by screening databases or libraries.
The
     compounds identified as binding partners or modulators of GPCR
CC
binding
     are useful for treating diseases in animals, and for control
insects that
     are harmful or cause injury to plants or animals. Diseases treated
     include infections (e.g. viral and human immunodeficiency virus,
CC
HIV),
CC
     cancer, pain, Parkinson's disease, hypotension, hypertension,
diabetes,
     obesity, atherosclerosis, thrombosis, stroke, renal failure,
     inflammation, rheumatoid arthritis, autoimmune disorders, and
psychotic
```

```
and neurological disorders (anxiety, schizophrenia, manic
depression,
    delirium, dementia, severe mental retardation, dyskinesias,
Huntington's
    disease or Tourette's syndrome). The nucleic acids can be used for
CC
CC
     genetic mapping, and producing the GPCRs. Anti-GPCR antibodies can
be
CC
     used in therapy, diagnostic assays and for modulating GPCR
activity.
XX
SQ
     Sequence
               14 AA;
                          96.3%; Score 52; DB 22; Length 14;
  Query Match
  Best Local Similarity 88.9%; Pred. No. 0.00088;
            8; Conservative 1; Mismatches 0; Indels
  Matches
Gaps
        1 FDDYGHLRF 9
Qу
          6 fddyghmrf 14
Db
SEQ ID NO: 22
RESULT
        1
AAU03215
    AAU03215 standard; Protein; 584 AA.
XX
AC
    AAU03215;
XX
DT
     12-SEP-2001 (first entry)
XX
    Fruit fly G protein coupled receptor, DmGPCR9.
DE
XX
KW
     Fruit fly; G protein coupled receptor; DmGPCR9;
KW
     human immunodeficiency virus; HIV; cancer; Parkinson's disease;
KW
     diabetes; obesity; atherosclerosis; thrombosis; stroke; renal
failure;
     inflammation; rheumatoid arthritis; autoimmune disorder;
KW
     neurological disorder; schizophrenia; manic depression; dementia;
KW
KW
     severe mental retardation; dyskinesia; Huntington's disease;
KW
    Tourette's syndrome.
XX
OS
    Drosophila melanogaster.
XX
PN
    WO200131005-A2.
XX
     03-MAY-2001.
PD
XX
     20-OCT-2000; 2000WO-US29002.
ΡF
XX
PR
     22-OCT-1999; 99US-0425676.
XX
     (PHAA ) PHARMACIA & UPJOHN CO.
PA
XX
```

```
PΤ
    Lowery DE, Smith VG, Kubiak TA, Larsen MJ;
XX
DR
    WPI; 2001-316333/33.
DR
    N-PSDB; AAS05894.
XX
    New Drosophila melanogaster GPCR nucleic acids and polypeptide
PΤ
useful
    for inducing an immune response, for identifying homologs and for
PΤ
    treating e.g. diabetes, obesity and manic depression -
PΤ
XX
PS
    Claim 29; Page 65; 110pp; English.
XX
    The sequence is a fruit fly G protein coupled receptor, DmGPCR9.
CC
    The proteins are useful for inducing an immune response against
CC
itself in
    a mammal. The nucleic acids are useful for identifying an animal
CC
homolog
    of DmGPCR, by screening databases or libraries. The compounds
identified
    as binding partners or modulators of GPCR binding are useful for
treating
    diseases in animals, and for control insects that are harmful or
CC
cause
    injury to plants or animals. Diseases treated include infections
CC
(e.g.
    viral and human immunodeficiency virus, HIV), cancer, pain,
CC
Parkinson's
    disease, hypotension, hypertension, diabetes, obesity,
atherosclerosis,
    thrombosis, stroke, renal failure, inflammation, rheumatoid
arthritis,
    autoimmune disorders, and psychotic and neurological disorders
(anxiety,
CC
    schizophrenia, manic depression, delirium, dementia, severe mental
    retardation, dyskinesias, Huntington's disease or Tourette's
CC
syndrome).
    The nucleic acids can be used for genetic mapping, and producing
    the GPCRs. Anti-GPCR antibodies can be used in therapy, diagnostic
CC
assays
    and for modulating GPCR activity.
CC
XX
SQ
               584 AA;
    Sequence
                         100.0%; Score 3000; DB 22;
 Query Match
                                                      Length 584;
 Best Local Similarity 100.0%; Pred. No. 1.8e-251;
 Matches 584; Conservative 0; Mismatches
                                                 0; Indels
                                                               0;
Gaps
       1 MFNYEEGDADQAAMAAAAYRALLDYYANAPSAAGHIVSLNVAPYNGTGNGGTVSLAGNA
QУ
60
          Db
       1 mfnyeegdadqaamaaaayralldyyanapsaaghivslnvapyngtgnggtvslagna
60
      61 TSSYGDDDRDGYMDTEPSDLVTELAFSLGTSSSPSPSSTPASSSSTSTGMPVWLIPSYSM
QУ
```

120

Db 120	61	
Qу 180	121	ILLFAVLGNLLVISTLVQNRRMRTITNVFLLNLAISDMLLGVLCMPVTLVGTLLRNFIFG
Db 180	121	
Qy 240	181	EFLCKLFQFSQAASVAVSSWTLVAISCERYYAICHPLRSRSWQTISHAYKIIGFIWLGGI
Db 240	181	
Qу 300	241	LCMTPIAVFSQLIPTSRPGYCKCREFWPDQGYELFYNILLDFLLVLPLLVLCVAYILIT
Db 300	241	
Qy 360	301	RTLYVGMAKDSGRILQQSLPVSATTAGGSAPNPGTSSSSNCILVLTATAVYNENSNNNNG
Db 360	301	
Qy 420	361	NSEGSAGGGSTNMATTTLTTRPTAPTVITTTTTTTTTTTTTAKTSSPSIRVHDAALRRSNEAK
Db 420	361	
Qy 480	421	TLESKKRVVKMLFVLVLEFFICWTPLYVINTMVMLIGPVVYEYVDYTAISFLQLLAYSSS
Db 480	421	
Qy 540	481	CCNPITYCFMNASFRRAFVDTFKGLPWRRGAGASGGVGGAAGGGLSASQAGAGPGAYASA
Db 540	481	
Qу	541	NTNISLNPGLAMGMGTWRSRSRHEFLNAVVTTNSAAAAVNSPQL 584
Db	541	ntnislnpglamgmgtwrsrsrheflnavvttnsaaaavnspql 584

.

9643744

```
PYRB LACLA
    PYRB LACLA
                 STANDARD;
                                PRT; 310 AA.
ID
AC
    Q9CF79;
DT
    16-OCT-2001 (Rel. 40, Created)
DT
    16-OCT-2001 (Rel. 40, Last sequence update)
    01-MAR-2002 (Rel. 41, Last annotation update)
DT
    Aspartate carbamoyltransferase (EC 2.1.3.2) (Aspartate
DE
DE
    transcarbamylase) (ATCase).
GN
    PYRB OR LL1602.
OS
    Lactococcus lactis (subsp. lactis) (Streptococcus lactis).
OC
    Bacteria; Firmicutes; Bacillus/Clostridium group; Streptococcaceae;
OC
    Lactococcus.
OX
    NCBI TaxID=1360;
RN
    [1]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=IL1403;
    MEDLINE=21235186; PubMed=11337471;
RA
    Bolotin A., Wincker P., Mauger S., Jaillon O., Malarme K.,
    Weissenbach J., Ehrlich S.D., Sorokin A.;
RA
    "The complete genome sequence of the lactic acid bacterium Lactococcus
RT
    lactis ssp. lactis IL1403.";
RT
    Genome Res. 11:731-753 (2001).
RI.
CC
    -!- CATALYTIC ACTIVITY: Carbamoyl phosphate + L-aspartate = phosphate
CC
        + N-carbamoyl-L-aspartate.
CC
    -!- PATHWAY: SECOND STEP IN PYRIMIDINE BIOSYNTHESIS.
CC
     -!- SIMILARITY: BELONGS TO THE ATCASES/OTCASES FAMILY.
CC
CC
    This SWISS-PROT entry is copyright. It is produced through a collaboration
CC
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
    the European Bioinformatics Institute. There are no restrictions on its
CC
CC
    use by non-profit institutions as long as its content is in no way
    modified and this statement is not removed. Usage by and for commercial
CC
CC
    entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
    or send an email to license@isb-sib.ch).
CC
    ______
DR
    EMBL; AE006390; AAK05700.1; -.
    InterPro; IPR002029; Carbmyltransf_asor.
DR
DR
    Pfam; PF00185; OTCace; 1.
    Pfam; PF02729; OTCace N; 1.
DR
    PRINTS; PR00100; AOTCASE.
DR
DR
    PROSITE; PS00097; CARBAMOYLTRANSFERASE; 1.
KW
    Pyrimidine biosynthesis; Transferase; Complete proteome.
SQ
    SEQUENCE 310 AA; 34558 MW; EEDE6B8EC6F00B94 CRC64;
 Query Match
                        72.2%; Score 39; DB 1; Length 310;
 Best Local Similarity 100.0%; Pred. No. 5;
 Matches
          6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qу
       1 FDDYGH 6
         Db
     201 FDDYGH 206
RESULT 12
RL5 SULSO
ID RL5 SULSO
                 STANDARD; PRT; 182 AA.
```

```
AC
    09UX93;
DT
    16-OCT-2001 (Rel. 40, Created)
    16-OCT-2001 (Rel. 40, Last sequence update)
DT
    01-MAR-2002 (Rel. 41, Last annotation update)
DT
DE
    50S ribosomal protein L5P.
GN
    RPL5P OR RPL5AB OR SSO0704 OR C10 026.
OS
    Sulfolobus solfataricus.
OC
    Archaea; Crenarchaeota; Sulfolobales; Sulfolobaceae; Sulfolobus.
OX
    NCBI TaxID=2287;
RN
    [1]
RP
    SEQUENCE FROM N.A.
    STRAIN=ATCC 35092 / DSM 1617 / P2;
RC
    MEDLINE=20165948; PubMed=10701121;
RX
    Charlebois R.L., Singh R.K., Chan-Weiher C.C.-Y., Allard G., Chow C.,
RA
    Confalonieri F., Curtis B., Duguet M., Erauso G., Faguy D.,
RA
    Gaasterland T., Garrett R.A., Gordon P., Jeffries A.C., Kozera C.,
RA
    Kushwaha N., Lafleur E., Medina N., Peng X., Penny S.L., She Q.,
RA
    St Jean A., van der Oost J., Young F., Zivanovic Y., Doolittle W.F.,
RA
    Ragan M.A., Sensen C.W.;
RA
     "Gene content and organization of a 281-kbp contig from the genome of
RT
     the extremely thermophilic archaeon, Sulfolobus solfataricus P2.";
RT
RL
    Genome 43:116-136(2000).
RN
     [2]
    SEQUENCE FROM N.A.
RP
RC
    STRAIN=ATCC 35092 / DSM 1617 / P2;
RX
    MEDLINE=21332296; PubMed=11427726;
    She Q., Singh R.K., Confalonieri F., Zivanovic Y., Allard G.,
RA
    Awayez M.J., Chan-Weiher C.C.-Y., Clausen I.G., Curtis B.A.,
RA
    De Moors A., Erauso G., Fletcher C., Gordon P.M.K.,
RA
    Heikamp-de Jong I., Jeffries A.C., Kozera C.J., Medina N., Peng X.,
RA
    Thi-Ngoc H.P., Redder P., Schenk M.E., Theriault C., Tolstrup N.,
RA
    Charlebois R.L., Doolittle W.F., Duguet M., Gaasterland T.,
RA
    Garrett R.A., Ragan M.A., Sensen C.W., Van der Oost J.;
RA
     "The complete genome of the crenarchaeon Sulfolobus solfataricus P2.";
RТ
     Proc. Natl. Acad. Sci. U.S.A. 98:7835-7840(2001).
RL
CC
     -!- SIMILARITY: BELONGS TO THE L5P FAMILY OF RIBOSOMAL PROTEINS.
CC
     ______
CC
    This SWISS-PROT entry is copyright. It is produced through a collaboration
CC
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
    the European Bioinformatics Institute. There are no restrictions on its
    use by non-profit institutions as long as its content is in no way
CC
CC
    modified and this statement is not removed. Usage by and for commercial
    entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
    or send an email to license@isb-sib.ch).
CC
CC
    EMBL; Y18930; CAB57599.1; -.
DR
    EMBL; AE006696; AAK41006.1; -.
DR
DR
    InterPro; IPR002132; Ribosomal L5.
DR
    Pfam; PF00281; Ribosomal L5; 1.
    Pfam; PF00673; Ribosomal L5 C; 1.
DR
DR
    ProDom; PD001076; Ribosomal_L5; 1.
DR
    PROSITE; PS00358; RIBOSOMAL L5; FALSE NEG.
KW
    Ribosomal protein; Complete proteome.
SQ
    SEQUENCE 182 AA; 20652 MW; 3070C4B01860D448 CRC64;
```

Best Local Similarity 66.7%; Pred. No. 4.4; Matches 6; Conservative 2; Mismatc

## 09693746 Results

SEQ ID NO: 22

### SUMMARIES

Result No	Score	Query Match	Length	DB	ID	Description
1	3000	100 0	584	22	AAU03215	Fruit fly G protei
2	1708	56 9	415	22	ABB62762	Drosophila melanog
3	1"08	56 9	415	22	AAU38944	Drosophila G-prote
4	1704	56 8	407	22	AAB86963	D. melanogaster pe
5	1242	41.4	749	22	ABB62718	Drosophila melanog
6	1242	41.4	749	22	AAU38943	Drosophila G-prote
7	970	32 3	201	22	AAB86962	D. melanogaster pe
8	964	32 1	205	22	ABB62772	Drosophila melanog
9	655.5	21 9	451	14	AAR40771	Sequence encoded b
10	655.5	21.9	452	22	AAB66619	Rat brain CCKB rec
11	654.5	21 8	430	14	AAR40772	Sequence encoded b
12	654.5	21 8	430	22	AAB66625	Guinea pig CCKA re
13	654.5	21 8	450	22	AAB66626	Guinea pig CCKA re
14	652.5	21 8	450	15	AAR53263	M. matalensis CCK
15	652.5	21 8	450	15	AAR59290	Mastomys gastrin r
16	647.5	21 6	428	18	AAW29102	Human peptide horm
17	647.5	21.6	428	22	AAB66630	Human CCK A recept
18	642	21.4	444	14	AAR38890	Sequence encoded b

```
RESULT
AAR40771
    AAR40771 standard; protein; 451 AA.
ID
XX
AC
     AAR40771;
XX
     07-FEB-1994 (first entry)
DT
XX
     Sequence encoded by the rat brain cholecystokinin (CCK) \ensuremath{\mathtt{B}}
DE
DE
     receptor cDNA clone.
XX
     Cholecystokinin receptor protein; CCK; gastrointestinal receptor.
КW
XX
os
     Balaenoptera acutorostrata
XX
FΗ
                     Location/Qualifiers
     Key
     Modified-site
FT
FT
                     /label= glycosylation site
                     /note= "see also AAs 30,36,255"
FT
FΤ
     Domain
                     57..80
FT
                     /label= transmembrane 1
FT
                     93. 116
     Domain
FT
                      /label= transmembrane II
FΤ
                     131. 150
     Domain
FT
                     /label= transmembrane III
FT
     Domain
                     173..192
FT
                     /label= transmembrane IV
FT
     Domain
                     219 .242
FT
                     /label= transmembrane V
FT
                     339 359
     Domain
                     /label= transmembrane VI
FT
FT
     Domain
                     374 381
FT
                     /label= transmembrane VII
XX
FΝ
     WO9316182-A.
XX
PD
     19-AUG-1993.
XX
ΡF
     28-JAN-1993; 93WO-US00466.
XX
```

```
07-FEB-1992; 92US-0831248.
01-APR-1992; 92US-0861769.
11-AUG-1992; 92US-0928033.
PR
PR
                92US-0937609.
PR
    02-SEP-1992;
XX
PΑ
    (USSH ) US DEPT HEALTH & HUMAN SERVICE.
XX
ΡI
ХХ
DR
    WPI; 1993-272886/34.
DR
    N-PSDB; AAQ47668.
XX
PΤ
    Isolated DNA molecule encoding cholecystokinin receptor protein -
    are purified to isolate cholecystokinin receptor clones and
PΤ
PΤ
    produce anti-cholecystokinin receptor antibodies
XX
PS
    Claim 19; Figure 2; 110pp; English.
XX
CC
    The rat brain CCK B receptor cDNA clone encodes a protein with
CC
    7 transmembrane domains, and homology with CCK A type receptor and
CC
    other G-protein receptor superfamily members. There are 4 potential
    sites of N-linked glycosylation, for serine phosphorylation
CC
CC
    (82,154,441), for disulphide bridges (127,205) and palmitoylation
CC
XX
SQ
    Sequence 451 AA;

        Query Match
        21.9%;
        Score 655.5;
        DB 14;
        Length 451;

        Best Local Similarity
        38.6%;
        Pred. No. 2.6e-48;

 Matches 164; Conservative 62; Mismatches 144; Indels 55; Gaps 11;
Q'/
      88 LGTSSSPSPSSTPASSSSTST---GMPVWLIPSYSMILLFAVLGNLLVISTLVQNRRMRT 144
         29 lnsssagnlscdpprirgtgtrelemai-ritlyaviflmsvggnvliivvlglsrrlrt 87
Db
     145 ITNVFLLNLAISDMLLGVLCMPVTLVGTLLRNFIFGEFLCKLFQFSQAASVAVSSWTLVA 204
Q;;
        Db
      88 vtnafllslavsdlllavacmpftllpnlmgtfifgtvickaisylmgvsvsvstlnlva 147
0.,
     205 ISCERYYAICHPLRSRSWQTISHAYKIIGFIWLGGILCMTPIAVFSQLIPTSRPGYCKCR 264
         Db
     148 ialerysaicrplqarvwqtrshaarvilatwllsgllmvpypvytmvqpvg-prvlqcm 206
     265 EFWPDQGYELFYNILLDFLLLVLPLLVLCVAYILITRTLYVGMAKD-----SGRILQ 316
Q:
          Db
     207 hrwpsarvqqtwsvllllllffipgvviavayglisrelylglhfdgendsetqsrarnq 266
     317 QSLPVSATTAGGSAPNPGTSSSSNCILVLTATAVYNENSNNNNGNSEGSAGGGSTNMATT 376
Qyr
     377 TLTTRPTAPTVITTTTTTTTTVTLAKTSSPSIRVHDAALRRSNEAKTLESKKRVVKMLFVLV 436
Qy
         Db
     306 rl-----emttlttptpgpvpgp-----rpnqakll-akkrvvrmllviv 344
Qy.
     437 LEFFICWTPLYVINTMVMLIGPVVYEYVDYTAISFLQLLAYSSSCCNPITYCFMNASFRR 496
          Db
     345 llfflcwlpvysvntwrafdgpgaqralsgapisfihllsyvsacvnplvycfmhrrfrq 404
     497 AFVDT 501
Qу
        i :||
     405 acldt 409
Issued:
```

Result Query
No. Score Match Length DB ID

Description

```
664.5 22 1 453 1 US-08-570-157-7
656 21 9 451 1 US-08-570-157-2
                                                           Sequence 7, Appli
                                                           Sequence 2, Appli
                     452 1 US-07-937-609-16
 3
     655 5
             21 9
                                                           Sequence 16, Appl
                              US-08-029-170-16
     655.5
             21 9
                     452 4
                                                           Sequence 16, Appl
                     430 1 US-07-937-609-23
             21.8
                                                           Sequence 23, Appl
     654 5
     654 5
             21 8
                     430 2 US-08-919-624-3
                                                           Sequence 3, Appli
                     430 4 US-08-029-170-23
450 1 US-07-937-609-24
     654 5
             21 8
                                                           Sequence 23, Appl
                                                           Sequence 24, Appl
 8
     654.5
             21.8
     654.5
             21.8
                     450 4 US-08-029-170-24
                                                           Sequence 24, Appl
 9
10
     648 5
             21.6
                     449 1
                              US-08-570-157-1
                                                           Sequence 1, Appli
                     428 1 US-08-570-157-5
11
     647.5
             21.6
                                                           Sequence 5, Appli
                     428 4 US-08-029-170-31
                                                           Sequence 31, Appl
12
     647 5
             21.6
     642
                     444 1 US-07-937-609-14
444 4 US-08-029-170-14
                                                           Sequence 14, Appl
             21.4
13
14
      642
             21.4
                                                           Sequence 14, Appl
                     443 1 US-08-570-157-6
     632 5
             21.1
                                                           Sequence 6, Appli
15
             20.9
                     453 1 US-07-937-609-26
16
     626.5
                                                           Sequence 26, Appl
     626 5
             20.9
                     453 4 US-08-029-170-26
448 1 US-08-570-157-3
                              US-08-029-170-26
17
                                                           Sequence 26, Appl
                                                           Sequence 3, Appli
      624
             20.8
1.8
19
     619.5
             20.6 447 1 US-07-937-609-29
                                                           Sequence 29, Appl
                     447 4
                                                           Sequence 29, Appl
20
     619 5
             20.6
                              US-08-029-170-29
                     453 1 US-07-937-609-27
     619 5
                                                           Sequence 27, Appl
21
             20.6
22
     613.5
             20.6 453 1 US-07-978-892A-5
                                                           Sequence 5, Appli
                     453 1 US-08-570-157-4
453 4 US-08-029-170-27
                                                           Sequence 4, Appli
23
     619.5
             20.6
24
     619.5
             20.6
                                                           Sequence 27, Appl
                   447 1 US-07-978-892A-6
     617.5
             20.6
                                                           Sequence 6, Appli
                   432 4 US-09-255-368-2
430 4 US-09-255-368-8
                                                           Sequence 2, Appli
     442
             14.7
26
27
     420.5
             14.0
                                                           Sequence 8, Appli
           13.8
28
     415.5
                     444 4 US-09-119-788-2
                                                           Sequence 2, Appli
     401
           13.4
                     425 4 US-09-479-128-2
29
                                                           Sequence 2, Appli
                     402 3 US-08-846-704-4
420 4 US-09-255-368-6
             13.3
30
       400
                                                           Sequence 4, Appli
           13 3
                                                           Sequence 6, Appli
31
      399
```

```
RESULT 1
US-08-570-157-7
; Sequence 7. Application US/08570157
; Patent No. 5750353
; GENERAL INFORMATION:
    APPLICANT: Kopin, Alan S.
     APPLICANT: Beinborn, Martin
    TITLE OF INVENTION: ASSAY FOR NON-PEPTIDE AGONISTS TO TITLE OF INVENTION: PEPTIDE HORMONE RECEPTORS
    NUMBER OF SEQUENCES: 23
     CORRESPONDENCE ADDRESS:
      ADDRESSEE: Fish & Richardson P.C.
      STREET: 225 Franklin Street
      CITY: Boston
STATE. MA
      COUNTRY: USA
      ZIP: 02110-2804
     COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
       COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/570,157
       FILING DATE: 11-DEC-1995
      CLASSIFICATION 435
    ATTORNEY/AGENT INFORMATION:
      NAME Clark, Paul T.
      FEGISTRATION NUMBER: 30,162
      PEFERENCE/DOCKET NUMBER 00398/109001
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 617/542-5070
      TELEFAX 617/542-8906
       TELEX: 200154
  INFORMATION FOR SEQ ID NO: 7:
    SEQUENCE CHAPACTERISTICS:
```

, LENGTH: 453 amino acids , TYPE: amino acid , STRANDEDNESS not relevant , TOPOLOGY: linear , MOLECULE TYPE: protein US-08-570-157-7

Best	Match 22 1%; Score 664.5; DB 1; Length 453; Local Similarity 33 5%; Pred No. 2.9e-45; es 176; Conservative 77; Mismatches 170; Indels 103; Gaps 16;
Q;′	13 AMAAAAAYRALLDYYANAPSAAGHIVSLNVAPYNGTGNGGTVSLAGNATSSYGDDDR 69 ::: :      :  :: ::       :
Db	6 SLSNISALHELLCRYSNLSGTLTWNLSSTNGTHNLTTANWPPWNLNCTPILDR 58
Q;;	70 DGYMDTEPSDLVTELAFSLGTSSSPSPSSTPASSSSTSTGMPVWL-IPSYSMILLFAVLG 128
Db	: : :     :   :    59LNLWVRIVMYSVIFLLSVFG 85
Q;·	29 NLLVISTLVQNRRMRTITNVFLLNLAISDMLLGVLCMPVTLVGTLLRNFIFGEFLCKLFQ 188
Db	:     : :        :  :  :::         :  :
Qyr	89 FSQAASVAVSSWTLVAISCERYYAICHPLRSRSWQTISHAYKIIGFIWLGGILCMTPIAV 248
Db	:   :  ::             :               ::   : ::     46 YFMGLSVSVSTFNLVAISIERYSAICNPLXSRVWQTRSHAYRVIAATWVLSSIIMIPYLV 205
Qyʻ	49 FSQLIPTSRPGYCKCREFWPDQGYELFYNILLDFLLVLPLLVLCVAYILITRTLY 304
Db	::: :
Qy	05 VGMAKDSGRILQQSLPVSATTAGGSAPNPGTSSSSNCILVLTATAVYNENSNNNNGNS 362
Db	: : : :             : :  65 RGIQFEMDLNKEAKAHKNGVSTPTTIPSGDEGDGCYIQVTKR 306
Qy	63 EGSAGGGSTNMATTTLTTRPTAPTVITTTTTTTTVTLAKTSSPSIRVHDAALRRSNEAKTL 422
Db	:     :
Qγ	23 ESKKRVVKMLFVLVLEFFICWTPLYVINTMVMLIGPVVYEYVDYTAISFLQLLAYSSSCC 482
Db	:    ::    :
Q <sub>2</sub> .	83 NPITYCFMNASFRRAFVDTFKGLPWRRGAGASGGVGGAAGGGLS 526
Db	

## SUMMARIES

Result No.	Score	Query Match	Length	DB	ID	Description
1	655.5	21.9	452	2	A46195	cholecystokinin B
2	654.5	21.8	430	2	I51898	cholecystokinin A
3	652.5	21 8	450	2	JQ1614	gastrin receptor -
4	647 5	21 6	428	2	JN0692	cholecystokinin ty
5	644.5	21.5	436	2	JC5599	cholecystokinin-A
6	642	21.4	444	2	A42685	cholecystokinın re
7	629.5	21.0	427	2	S50150	gastric CCK-A rece
8	625	20.8	452	2	JC2459	gastrin/cholecysto
9	619.5	20.6	453	2	S32817	gastrin receptor -
10	617.5	20.6	447	2	A47430	gastrin/cholecysto
11	459.5	15.3	381	2	S48049	cholecystokinin B
12	427.5	14.2	397	2	T25910	hypothetical prote
13	420.5	14 0	449	2	A41738	neuropeptide Y rec
14	382.5	12 8	465	1	JQ1517	neurokinin 3 recep
15	372 5	12 4	452	2	A34916	neurokinin 3 recep
16	370.5	12 3	407	2	S23510	neurokinin 1 recep
17	367 5	12.2	440	2	A44081	kappa-type opioid

```
RESULT
A46195
cholecystokinin B receptor subtype - rat
C; Species: Rattus norvegicus (Norway rat)
C,Date: 21-Sep-1993 #sequence revision 18-Nov-1994 #text change 20-Apr-2000
C.Accession: A46195
R; Wank, S.A.; Pisegna, J.R.; de Weerth, A.
Proc. Natl. Acad. Sci. U.S.A. 89, 8691-8695, 1992
A:Title: Brain and gastrointestinal cholecystokinin receptor family structure and
functional expression
A; Reference number · A46195; MUID: 92409582
A; Accession: A46195
A; Status: preliminary
A; Molecule type: nucleic acid
A; Residues: 1-452 <WAN>
A;Cross-references. GB:M99418; NID:g203459; PIDN:AAA40925.1; PID:g203460
A; Experimental source brain
A; Note: sequence extracted from NCBI backbone (NCBIN:114083, NCBIP:114084)
C; Superfamily: neurokinin 1 receptor
C; Keywords: G protein-coupled receptor; transmembrane protein
 Query Match 21.9%; Score 655.5; DB 2; Length 452; Best Local Similarity 38.6%; Pred. No. 2.5e-38;
 Matches 164; Conservative 62; Mismatches 144; Indels 55; Gaps 11;
      88 LGTSSSPSPSSTPASSSSTST---GMPVWLIPSYSMILLFAVLGNLLVISTLVQNRRMRT 144
        29 LNSSSAGNLSCDPPRIRGTGTRELEMAI-RITLYAVIFLMSVGGNVLIIVVLGLSRRLRT 87
Db
01.
     145 ITNVFLLNLAISDMLLGVLCMPVTLVGTLLRNFIFGEFLCKLFQFSQAASVAVSSWTLVA 204
        88 VTNAFLLSLAVSDLLLAVACMPFTLLPNLMGTFIFGTVICKAISYLMGVSVSVSTLNLVA 147
Db
Qy
     205 ISCERYYAICHPLRSRSWQTISHAYKIIGFIWLGGILCMTPIAVFSQLIPTSRPGYCKCR 264
         148 IALERYSAICRPLOARVWOTRSHAARVILATWLLSGLLMVPYPVYTMVOPVG-PRVLOCM 206
Db
01.7
     265 EFWPDQGYELFYNILLDFLLLVLPLLVLCVAYILITRTLYVGMAKD-----SGRILQ 316
               Db
     207 HRWPSARVQQTWSVLLLLLLFFIPGVVIAVAYGLISRELYLGLHFDGENDSETQSRARNQ 266
Qyr
     317 QSLPVSATTAGGSAPNPGTSSSSNCILVLTATAVYNENSNNNNGNSEGSAGGGSTNMATT 376
     Db
0::
     377 TLTTRPTAPTVITTTTTTTTTLAKTSSPSIRVHDAALRRSNEAKTLESKKRVVKMLFVLV 436
                   306 RL------EMTTLTTPTPGPVPGP-----RPNQAKLL-AKKRVVRMLLVIV 344
Db
Qy.
     437 LEFFICWTPLYVINTMVMLIGPVVYEYVDYTAISFLQLLAYSSSCCNPITYCFMNASFRR 496
        345 LLFFLCWLPVYSVNTWPAFDGPGAQRALSGAPISFIHLLSYVSACVNPLVYCFMHRRFRQ 404
Db
     497 AFVDT 501
Qy
        | :||
Db
     405 ACLDT 409
RESULT 2
cholecystokinin A receptor - guinea pig
C, Species Cavia porcellus (quinea pig)
C,Date: 04-Sep-1997 #sequence_revision 04-Sep-1997 #text_change 20-Apr-2000
C,Accession: I51898
R,De Weerth, A.; Pisegna, J.R.; Wank, S.A.
Am. J. Physiol. 265, G1116-G1121, 1993
```

```
A; Title: Guinea pig gallbladder and pancreas possess identical CCK-A receptor subtypes:
receptor cloning and expression.
A; Reference number: I51898, MUID: 94106629
A; Accession: I51898
A.Status: preliminary; translated from GB/EMBL/DDBJ
A.Molecule type: mRNA
A, Residues: 1-430 < RES>
A.Cross-references: GB·S68242; NID:g544723; PIDN:AAB29504.1, PID g544724
C, Superfamily: neurokinin 1 receptor
 Query Match 21 8%; Score 654.5; DB 2; Length 430; Best Local Similarity 35 0%; Pred. No. 2.7e-38;
                           72; Mismatches 168; Indels 67, Gaps 10;
 Matches 165; Conservative
      83 ELAF---SLGTSSSPSPSSTPASSSSTSTGMPVWLIPSYSMILLFAVLGNLLVISTLVQN 139
01.
         19 ELGFENETLFCLDRPRPS------KEWQPAVQILLYSLIFLLSVLGNTLVITVLIRN 69
Qy.
     140 RRMRTITNVFLLNLAISDMLLGVLCMPVTLVGTLLRNFIFGEFLCKLFQFSQAASVAVSS 199
         70 KRMRTVTNIFLLSLAVSDLMLCLFCMPFNLIPSLLKDFIFGSAVCKTTTYFMGTSVSVST 129
Db
     200 WTLVAISCERYYAICHPLRSRSWQTISHAYKIIGFIWLGGILCMTPIAVFSQLIPTSRPG 259
Qv.
         130 FNLVAISLERYGAICKPLQSRVWQTKSHALKVIAATWCLSFTIMTPYPIYSNLVPFTKNN 189
Qy/
     260 Y---CKCREFWPDQGYELFYNILLDFLLLVLPLLVLCVAYILITRTLYVGMAKDSGRILQ 316
             190 NOTGNMCRFLLPNDVMQQTWHTFLLLILFLIPGIVMMVAYGLISLELYQGIKFDA--IQK 247
Db
     317 QSLPVSATTAGGSAPNPGTSSSSNCILVLTATAVYNENSNNNNGNSEGSAGGGSTNMATT 376
017
     | | | | | | | | | | | | 248 KSAKERKTSTGSSGP---MEDSDGC------YLQKSRH-----PR 278
Db
Qy
     377 TLTTRPTAPTVITTTTTTTTTLAKTSSPSIRVHDAALRRSNEAKTLESKKRVVKMLFVLV 436
                           279 FLELROLSP------SSSGSNRIN--RIRSSSSTANLMAKERVIRMLIVIV 321
Db
     437 LEFFICWTPLYVINTMVMLIGPVVYEYVDYTAISFLQLLAYSSSCCNPITYCFMNASFRR 496
Q٧
                             :: | | | | : | | : | ' | | | | | | | |
         322 VLFFLCWMPIFSANAWRAYDTVSAERHLSGTPISFILLLSYTSSCVNPIIYCFMNKRFRL 381
Db
QУ
     497 AFVDTFKGLPWRRGAGASGGVGGAAGGGLSASQAGAGPGAYASANTNISLNP 548
                 : | | | | : : : | |
         1: 11
     382 GFMATFPCCP----NPGTPGVRGEMGEEEEGRTTGASLSRYSYSHMSTSAPP 429
Db
```

#### SUMMARIES

Result		% Query				
No.	Score		Length	DB	ID	Description
1	667.5	22 2	453	1	CCEP_XENLA	P70031 xenopus lae
2	655.5	21 9	452	1	GASP_RAT	P30553 rattus norv
3	654.5	21.8	430	1	CCHR_CAVPO	Q63931 cavia porce
4	652 5	21.8	450	1	GASE PRANA	P30796 praomys nat
5	647 5	21.6	428	1	CCER_HUMAN	P32238 homo sapien
6	646.5	21.6	453	1	GASPMOUSE	P56481 mus musculu
7	644.5	21.5	436	1	CCKR MOUSE	008786 mus musculu
8	642	21.4	444	1	CCER_RAT	P30551 rattus norv
9	632.5	21.1	427	1	CCKR_RABIT	097772 oryctolagus
10	625	20 8	452	1	GASE_RABIT	P46627 oryctolagus
11	619 5	20 6	453	1	GASR_CANFA	P30552 canis famil
12	617.5	20.6	447	1	GASR HUMAN	P32239 homo sapien
13	612	20 4	454	1	GASR_BOVIN	P79266 bos taurus
14	442	14 7	432	1	NFF1_RAT	Q9ep86 rattus norv

```
RESULT 1
CCKR XENLA
    CCKR XENLA
                    STANDARD;
                                   PRT; 453 AA.
ID
    P70031:
     01-NOV-1997 (Rel. 35, Created)
DT
    01-NOV-1997 (Pel. 35, Last sequence update)
T)T
    15-JUL-1998 (Rel. 36, Last annotation update)
DT
    Cholecystokinin receptor (CCK-XLR).
DE
    Xenopus laevis (African clawed frog).
CiS
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC.
     Amphibia; Batrachia; Anura; Mesobatrachia; Pipoidea; Pipidae;
OC.
OC.
     Xenopodinae; Xenopus.
OX
    NCBI_TaxID=8355;
PN
    [1]
P.P
     SEQUENCE FROM N.A.
RC
     TISSUE=Brain:
RX
    MEDLINE=96319796; PubMed=8700154;
     Schmitz F., Pratt D S., Wu M.-J., Kolakowski L F. Jr., Beinborn M.,
F:A
PΑ
     Kopin A.S.;
PT
     "Identification of cholecystokinin-B/gastrin receptor domains that
     confer high gastrin affinity: utilization of a novel Xenopus laevis
ŖТ
     cholecystokinin receptor.";
RΤ
     Mol. Pharmacol. 50:436-441(1996).
P.L
     -!- FUNCTION. RECEPTOR FOR CHOLECYSTOKININ THIS RECEPTOR MEDIATES ITS
CC
         ACTION BY ASSOCIATION WITH G PROTEINS THAT ACTIVATE A
CC
         PHOSPHATIDYLINOSITOL-CALCIUM SECOND MESSENGER SYSTEM. HAS HIGH
CC
         AFFINITY FOR CCK-8 AND LOW AFFINITIES FOR GASTRIN-17-I, CCK-4, AND
CC
CC
         UNSULFATED CCK-8.
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein.
     -!- TISSUE SPECIFICITY: BRAIN AND STOMACH.
CC
     -!- SIMILARITY: BELONGS TO FAMILY 1 OF G-PROTEIN COUPLED RECEPTORS.
CC
        HAS EQUAL SIMILARITY TO TYPE A AND B CHOLECYSTOKININ MAMMALIAN
CC
CC
        RECEPTORS.
CC
     This SWISS-PROT entry is copyright. It is produced through a collaboration
CC
CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
     the European Bioinformatics Institute. There are no restrictions on its
     use by non-profit institutions as long as its content is in no way
CC
CC
     modified and this statement is not removed. Usage by and for commercial
CC
     entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
     or send an email to license@isb-sib.ch).
CC
    EMBL; U49258; AAB09052.1; -.
DR
DR
    GCRDb; GCR 0930; -.
     GCRDb; GCR 1953; -.
DR
DR
    InterPro: IPR000276: GPCR Rhodpsn.
DR
     Pfam; PF00001; 7tm_1; 1.
     PRINTS; PR00237; GPCRRHODOPSN
DR
    PROSITE; PS00237; G_PROTEIN_RECEP_F1_1; 1.
DR
     PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
DR
     G-protein coupled receptor; Transmembrane; Glycoprotein;
EW
EW
    Lipoprotein; Palmitate.
FT
     DOMAIN
                  1
                        64
                                  EXTRACELLULAR (POTENTIAL).
                        94
FT
     TRANSMEM
                  65
                                  1 (POTENTIAL)
FT
                 95
                       104
                                  CYTOPLASMIC (POTENTIAL).
     DOMAIN
FT
     TRANSMEM
                 105
                        131
                                  2 (POTENTIAL)
FT
    DOMAIN
                 132
                        142
                                  EXTRACELLULAR (POTENTIAL).
                                  3 (POTENTIAL)
FT
     TRANSMEM
                 143
                        164
                                  CYTOPLASMIC (POTENTIAL) .
FT
     DOMAIN
                 165
                        184
FT
     TRANSMEM
                 185
                        205
                                  4 (POTENTIAL)
FT
     DOMAIN
                 206
                        237
                                  EXTRACELLULAR (POTENTIAL).
FT
     TRANSMEM
                 238
                        261
                                  5 (POTENTIAL)
FT
     DOMAIN
                 262
                        343
                                  CYTOPLASMIC (POTENTIAL).
FT
     TRANSMEM
                 344
                        364
                                  6 (POTENTIAL)
                        379
                                  EXTRACELLULAR (POTENTIAL).
FT
    DOMAIN
                 365
FT
                 380
                        403
                                  7 (POTENTIAL)
     TRANSMEM
FT
                                  CYTOPLASMIC (POTENTIAL).
    DOMAIN
                 404
                        453
FT
    DISULFID
                 141
                        223
                                  BY SIMILARITY
FT
                                  PALMITATE (BY SIMILARITY).
     LIPID
                 401
                        401
                                 N-LINKED (GLCNAC . .) (POTENTIAL) .
N-LINKED (GLCNAC . .) (POTENTIAL) .
    CARBOHYD
                 9
FT
                        9
FT
     CARBOHYD
                 22
                      2.2
```

FT	CARBOHYD 30 30 N-LINKED (GLCNAC) (POTENTIAL). CARBOHYD 35 35 N-LINHED (GLCNAC) (POTENTIAL). CARBOHYD 39 39 N-LINHED (GLCNAC) (POTENTIAL). SEQUENCE 453 AA; 51157 MW; 06217927B7482678 CRC64;
Que Bes Mat	ery Match 22 2%, Score 667.5; DB 1, Length 453; st Local Similarity 33 5%. Pred No. 6.4e-34; tches 176; Conservative 78, Mismatches 169. Indels 103; Gaps 16;
QУ	13 AMAAAAAYRALLDYYANAPSAAGHIVSLNVAPYNGTGNGGTVSLAGNATSSYGDDDR 69
Db	6 SLSNISALHELLCRYSNLSGT LTWNLSSTNGTHNLTTANWPPWNLNCTPILDR 58
Qу	70 DGYMDTEPSDLVTELAFSLGTSSSPSPSSTPASSSSTSTGMPVWL-IPSYSMILLFAVLG 128
Db	
QУ	129 NLLVISTLVQNRRMRTITNVFLLNLAISDMLLGVLCMPVTLVGTLLRNFIFGEFLCKLFQ 188
Db	86 NTLIIIVLVMNKRLRTITNSFLLSLALSDLMVAVLCMPFTLIPNLMENFIFGEVICRAAA 145
QУ	189 FSQAASVAVSSWTLVAISCERYYAICHPLPSRSWQTISHAYKIIGFIWLGGILCMTPIAV 248
Db	146 YFMGLSVSVSTFNLVAISIERYSAICNPLKSRVWQTRSHAYRVIAATWVLSSIIMIPYLV 205
QУ	249 FSQLIPTSRPGYCKCREFWPDQGYELFYNILLDFLLVLPLLVLCVAYILITRTLY 304
Db	:::::         ::           :   :   :
QУ	305 VGMAKDSGRILQQSLPVSATTAGGSAPNPGTSSSSNCILVLTATAVYNENSNNNNGNS 362
Db	: : :           265 RGIQFEMDLNKEAKAHKNGVSTPTTIPSGDEGDGCYIQVTKR 306
Q٧	363 EGSAGGGSTNMATTTLTTRPTAPTVITTTTTTTTTTTTTTAKTSSPSIRVHDAALRRSNEAKTL 422
Db	:     :     ::      307
QУ	423 ESKKRVVKMLFVLVLEFFICWTPLYVINTMVMLIGPVVYEYVDYTAISFLQLLAYSSSCC 482
Db	:    ::   :       ::    ::      : : :   336 MAKKRVIRMLIVIVAMFFICWMPIFVANTWKAFDELSAFNTLTGAPISFIHLLSYTSACV 395
QУ	483 NPITYCFMNASFRRAFVDTFKGL- PWRRGAGASGGVGGAAGGGLS 526
Db	

SEQ ID NO: 157

8						
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	54	100.0	9	22	AAU03346	Fruit fly G protei
2	54	100 0	9	22	AAU03347	Fruit fly G protei
3	54	100.0	9	22	AAU03351	Fruit fly G protei
4	52	96 3	14	22	AAU03353	Fruit fly G protei
5	52	96.3	128	22	ABB66665	Drosophila melanog
б	4.8	88.9	9	22	AAU03897	G protein-coupled
7	40	74.1	7	22	AAU03354	G protein coupled
8	40	74.1	590	22	AAB84261	Amino acid sequenc
9	40	74.1	640	22	ABG16509	Novel human diagno
10	40	74 1	836	19	AAW85017	Grk5-green floures
11	40	74 1	842	19	AAW85008	Grk5-green floures
12	3,9	72 2	227	22	AAU31534	Novel human secret
13	39	72 2	255	22	ABB71872	Drosophila melanog
14	39	72 2	345	22	AAU33607	Pseudomonas aerugi
15	3.8	70.4	281	22	ABB59929	Drosophila melanog
16	37	68 5	89	22	AAM16451	Peptide #2885 enco

17	37	68.5	145	22	AAM25650	Human protein sequ
18	37	68.5	431	20	AAY59728	Human normal ovari
19	37	68.5	855	21	AAB54359	Human pancreatic c
20	37	68.5	1090	22	AAB94737	Human protein sequ
21	37	68 5	1144	21	AAB02007	Type III adenylyl
22	36	66 7	149	22	AAM79774	Human protein SEQ
23	36	66.7	244	21	AAG07361	Arabidopsis thalia
24	36	66 7	244	21	AAG61263	Arabidopsis thalia
25	36	66 7	251	21	AAG07360	Arabidopsis thalia
26	36	66 7	251	21	AAG61262	Arabidopsis thalia
27	36	66 7	253	22	AAB86351	A. thaliana allene
28	36	66 7	254	22	AAB86350	A. thaliana allene
29	36	66 7	258	21	AAG07359	Arabidopsis thalia
30	36	66 7	258	21	AAG61261	Arabidopsis thalia
31	36	66 7	301	22	AAM78790	Human protein SEQ

#### SUMMARIES

Result Query Score Match Length DB ID Description \_\_\_\_ 74.1 590 1 US-08-221-817-14 74.1 590 1 US-08-454-439-14 74.1 590 4 US-08-464-954A-5 40 Sequence 14, Appl 40 Sequence 14, Appl Sequence 5, Appli 3 40 40 74.1 590 5 PCT-US94-10487-14 Sequence 14, Appl 1144 3 US-08-726-214-6 576 1 US-08-221-817-13 5 37 68.5 Sequence 6, Appli 6 35 64.8 Sequence 13, Appl 35 64.8 576 1 US-08-221-817-22 Sequence 22, Appl 576 1 US-08-454-439-13 576 1 US-08-454-439-22 35 8 64 8 Sequence 13, Appl 9 35 64.8 Sequence 22, Appl 576 4 US-08-464-954A-6 10 35 64.8 Sequence 6, Appli 11 35 64.8 576 5 PCT-US94-10487-13 Sequence 13, Appl 12 35 64.8 576 5 PCT-US94-10487-22 Sequence 22, Appl 632 1 US-08-221-817-11 Sequence 11, Appl 13 35 64.8 14 35 64.8 632 1 US-08-454-439-11 Sequence 11, Appl 15 35 64.8 632 5 PCT-US94-10487-11 Sequence 11, Appl 35 64 8 688 1 US-08-221-817-19 16 Sequence 19, Appl 17 35 64 8 688 1 US-08-454-439-19 Sequence 19,

```
RESULT 1
US-08-221-817-14
; Sequence 14, Application US/08221817
; Patent No. 5532151
  GENERAL INFORMATION:
    APPLICANT: Chantry, David
    APPLICANT: Gray, Patrick W.
    APPLICANT: Hoekstra, Merle F.
    TITLE OF INVENTION: A No. 5532151el G Protein-Coupled Receptor TITLE OF INVENTION: Kinase GRK6
    NUMBER OF SEQUENCES: 24
     CORRESPONDENCE ADDRESS:
      ADDRESSEE Marshall, O'Toole, Gerstein, Murray &
      ADDRESSEE Borun
      STREET: 6300 Sears Tower, 233 South Wacker Drive
      CITY: Chicago
      STATE: Illinois
      COUNTRY: USA
      ZIP: 60606
     COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Pelease #1.0, Version #1.25
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER US/08/221,817
      FILING DATE
      CLASSIFICATION: 435
```

PRIOR APPLICATION DATA:

APPLICATION NUMBER 08/123,932

```
FILING DATE: 17 SEP 1993
ATTORNEY/AGENT INFORMATION:
      NAME: No. 5532151and, Greta E.
       REGISTRATION NUMBER: 35,302
       REFERENCE/DOCKET NUMBER: 31981
     TELECOMMUNICATION INFORMATION
       TELEPHONE (312) 474-6300
       TELEFAX (312: 474-0448
      TELEX: 25-3856
   INFORMATION FOR SEQ ID NO: 14:
     SEQUENCE CHARACTERISTICS:
      LENGTH: 590 amino acids
       TYPE. amino acid
       TOPOLOGY linear
     MOLECULE TYPE: protein
US-08-221-817-14
 Query Match
                          74.1%; Score 40; DB 1; Length 590;
 Best Local Similarity 85.7%; Pred. No. 10;
  Matches 6, Conservative 1; Mismatches
                                                   0; Indels
                                                                 0; Gaps
        2 DDYGHLR 8
          1 111:1
      320 DDYGHIR 326
RESULT 2
US-08-454-439-14
: Sequence 14, Application US/08454439
. Patent No. 5591618
 GENERAL INFORMATION:
     APPLICANT: Chantry, David
     APPLICANT: Gray, Patrick W. APPLICANT: Hoekstra, Merle F.
     TITLE OF INVENTION. A No. 5591618el G Protein-Coupled Receptor
    TITLE OF INVENTION Kinase GRK6
NUMBER OF SEQUENCES: 24
     CORRESPONDENCE ADDRESS
      ADDRESSEE: Marshall, O'Toole, Gerstein, Murray & ADDRESSEE: Borun
       STREET: 6300 Sears Tower, 233 South Wacker Drive
      CITY: Chicago
STATE: Illinois
       COUNTRY: USA
       ZIP: 60606
     COMPUTER READABLE FORM
       MEDIUM TYPE. Floppy disk
       COMPUTER: IBM PC compatible
       OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.25
     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/08/454,439
      FILING DATE: 30-MAY-1995
      CLASSIFICATION 435
     PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US 08/221,817
       FILING DATE: 31-MAR-1994
       APPLICATION NUMBER: 08/123,932
      FILING DATE: 17 SEP 1993
       CLASSIFICATION: 435
     ATTORNEY/AGENT INFORMATION:
      NAME: No. 5591618and, Greta E.
       REGISTRATION NUMBER: 35,302
      REFERENCE/DOCKET NUMBER: 31981
     TELECOMMUNICATION INFORMATION.
      TELEPHONE: (312) 474-6300
       TELEFAX: (312) 474-0448
       TELEX 25-3856
  INFORMATION FOR SEQ ID NO: 14:
    SEQUENCE CHARACTERISTICS:
```

```
LENGTH 590 amino acids
       TYPE: amino acid
       TOPOLOGY: linear
    MOLECULE TYPE · protein
US-08-454-439-14
 Query Match 74.1%; Score 40; DB 1; Length 590; Best Local Similarity 85.7%; Pred. No. 10; Matches 6; Conservative 1; Mismatches 0; Indels
        2 DDYGHLR 8
          320 DDYGHIR 326
Dh
Result
                Query
        Score Match Length DB ID
                                                             Description
  No.
  ______
           52 96.3
52 96.3
                         11 2 A60656
14 2 A56632
                                                              perisulfakinin - A
     1
                                                              neosulfakinin-II -
     2
           52 96 3 128 2 A31101
                                                              drosulfakinin prec
           52 96 3 140 2 S66610
48 88 9 11 1 GMROL
                                                              sulfakinin - blueb
     4
     5
                                                              leucosulfakinin -
                         10 1 GMROL2
           46 85.2
                                                             leucosulfakinin-II
     6
           46 85.2
40 74.1
                         10 2 B60656
35 2 B48682
     7
                                                              leucosulfakinin II
           40 74.1 35 2 B48682
40 74.1 590 1 A54372
     8
                                                             G protein-coupled
     9
                                                             G protein-coupled
           40 74.1 590 2 A48277
39 72.2 310 2 B86825
                                                             G protein-coupled
    10
                                                              aspartate carbamoy
    11
           39 72.2 310 2 B86825
39 72.2 345 2 B83371
                                                              conserved hypothet
    1.2
           39 72.2 419 2 S72325
39 72.2 471 2 B97611
39 72.2 471 2 AF2833
    13
                                                             glucan 1,3-beta-gl
                                                              UDP-N-acetylmurama
    14
                                                              UDP-N-acetylmurama
    15
RESULT
A60656
perisulfakinin - American cockroach
C; Species: Periplaneta americana (American cockroach)
C;Date: 14-May-1993 #sequence_revision 14-May-1993 #text_change 11-Jul-1997
C; Accession: A60656
R: Veenstra, J.A.
Neuropeptides 14, 145-149, 1989
A; Title: Isolation and structure of two gastrin/CCK-like neuropeptides from the American
cockroach homologous to the leucosulfakinins.
A; Reference number: A60656; MUID: 90137190
A; Accession: A60656
A; Molecule type: protein
A Residues: 1-11 <VEE>
```

```
C;Comment: This neuropeptide stimulates hindgut contractions.
C,Keywords: amidated carboxyl end; neuropeptide; sulfoprotein
F,6/Binding site: sulfate (Tyr) (covalent) #status experimental
F,11/Modified site. amidated carboxyl end (Phe) #status experimental

Query Match 96.3%; Score 52; DB 2; Length 11;
Best Local Similarity 88.9%; Pred. No. 0.0013;
Matches 8; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 1 FDDYGHLRF 9
| | | | | | | | |
Db 3 FDDYGHMRF 11

RESULT 2
A56632
neosulfakinin-II - flesh fly (Sarcophaga bullata)
```

N; Alternate names: Neb-SK-II

N, Contains: neosulfakinin-I (Neb-SK-I) C. Species: Sarcophaga bullata C;Date: 21-Jul-1995 #sequence\_revision 21-Jul-1995 #text\_change 20-Jun-2000 C; Accession: A56632 R,Fonagy, A.; Schoofs, L.: Proost, P; Van Damme, J.; De Loof, A Comp. Biochem Physiol. C 103, 135-142, 1992 A, Title: Isolation and primary structure of two sulfakinin-like peptides from the fleshfly, Neobellieria bullata. A, Reference number: A56632; MUID: 93083101 A, Accession: A56632 A, Molecule type: protein A; Residues: 1-14 < FON> A, Experimental source: heads A, Note: sequence extracted from NCBI backbone (NCBIP:120391) C; Keywords: amidated carboxyl end; neuropeptide; sulfoprotein F;1-14/Product: neosulfakinin-II #status experimental <NSK2> F,6-14/Product: neosulfakının-I #status experimental <NSK1> F,9/Binding site: sulfate (Tyr) (covalent) #status predicted F,14/Modified site: amidated carboxyl end (Phe) #status experimental Query Match 96.3%; Score 52; DB 2; Length 14; Best Local Similarity 88.9%; Pred. No. 0.0016; Matches 8; Conservative 1; Mismatches 0; Indels 0; Gaps 0; 1 FDDYGHLRF 9  $|\cdot|\cdot|\cdot|\cdot|$ 6 FDDYGHMRF 14 SUMMARIES

		8				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	52	96.3	9	1	NSK1_SARBU	P41492 sarcophaga
2	52	96.3	11	1	LSKP_PERAM	P36885 periplaneta
3	52	96.3	14	1	NSK2_SARBU	P41493 sarcophaga
4	52	96.3	128	1	DSK_DROME	P09040 drosophila
5	48	88.9	11	1	LSK1_LEUMA	P04428 leucophaea
6	46	85.2	10	1	LSF.2_LEUMA	P09039 leucophaea
7	46	85.2	12	1	LOSK_LOCMI	P47733 locusta mig
8	40	74.1	590	1	GRK5_BOVIN	P43249 bos taurus
9	40	74.1	590	1	GRES_HUMAN	P34947 homo sapien
10	40	74.1	590	1	GRK5_RAT	Q62833 rattus norv
11	39	72.2	310	1	PYPB_LACLA	Q9cf79 lactococcus
12	38	70.4	182	1	RL5_SULSO	Q9ux93 sulfolobus
13	38	70.4	463	1	FLGE_TREPH	Q56326 treponema p
14	37	68.5	190	1	RL5_METJA	P54040 methanococc
15	37	68.5	1144	1	CYA3_HUMAN	060266 homo sapien
16	37	68.5	1144	1	CYA3_RAT	P21932 rattus norv
17	36	66.7	200	1	YCLP_XANCP	P22264 xanthomonas
18	36	66.7	214	1	VC01_VARV	P33859 variola vir
19	36	66.7	224	1	VC01_VACCC	P21036 vaccinia vi
20	36	66.7	229	1	VC01_VACCV	P17368 vaccinia vi
21	36	66.7	454	1	MUP.C_AQUAE	067373 aquifex aeo
22	35	64.8	576	1	GRE6_HUMAN	P43250 homo sapien
23	35	64.8	576	1	GRK6_MOUSE	070293 mus musculu
24	35	64.8	576	1	GRE6_RAT	P97711 rattus norv
25	35	64.8	642	1	YQR1_CAEEL	Q09537 caenorhabdi

```
RESULT 1
NSK1_SARBU

ID NSK1_SARBU STANDARD; PET; 9 AA.
AC P41492;
DT 01-NOV-1995 (Rel. 32, Created)
DT 01-NOV-1995 (Rel. 32, Last sequence update)
```

```
DT
    01-FEB-1996 (Rel. 33, Last annotation update)
     Neosulfakinin-I (NEB-SK-I)
     Sarcophaga bullata (Grey flesh fly) (Neobellieria bullata).
CS
C:C
     Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
C.C
     Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
CC
     Oestroidea; Sarcophagidae; Sarcophaga
C:X
     NCBI_TaxID=7385;
F.N
     [1]
     SEQUENCE.
FP
F.C
     TISSUE=Head:
     MEDLINE=93083101; PubMed=1360367.
RХ
P.A
     Fonagy A., Schoofs L., Proost P., van Damme J., de Loof A.;
     "Isolation and primary structure of two sulfakinin-like peptides from
FT
PT
     the fleshfly, Neobellieria bullata.";
F.L
     Comp. Biochem. Physiol. 103C:135-142(1992).
     -!- FUNCTION: MYOTROPIC PEPTIDE.
CC
CC
     -!- SIMILARITY: BELONGS TO THE GASTRIN/CHOLECYSTOKININ FAMILY.
DR
     InterPro; IPR001651; Gastrin.
     PROSITE; PS00259; GASTRIN; 1.
Ľ.R
F.W
     Neuropeptide; Amidation; Sulfation.
               4 4 SULFATION (POTENTIAL).
9 9 AMIDATION (POTENTIAL).
FΤ
     MOD RES
     MOD RES
FT
     SEQUENCE 9 AA; 1187 MW; 8B0A0691E86B5AAA CRC64;
 Query Match
                           96.3%; Score 52, DB 1; Length 9;
 Best Local Similarity 88.9%; Pred. No. 1e+05;
Matches 8; Conservative 1; Mismatches 0; Indels
                                                                  0; Gaps
        1 FDDYGHLRF 9
QУ
          11111:11
        1 FDDYGHMRF 9
Db
                                     SUMMARIES
```

		*5				
Result		Query				
No	Score	Match	Length	DB	ID	Description
1	40	74.1	590	11	070292	070292 mus musculu
2	40	74.1	590	11	070297	070297 mus musculu
3	39	72.2	255	5	Q9VIT0	Q9vıt0 drosophila
4	39	72.2	310	2	Q9L4N6	Q914n6 lactococcus
5	39	72.2	345	16	Q9I1S0	Q9ils0 pseudomonas
6	39	72.2	419	3	Q12539	Q12539 agaricus bi
7	39	72.2	420	3	Q9C1A8	Q9cla8 gibberella
8	39	72.2	420	3	Q9C1B5	Q9c1b5 fusarium sp
9	39	72.2	420	3	Q96V36	Q96v36 gibberella
10	39	72.2	466	16	Q98KB4	Q98kb4 rhizobium l
11	39	72.2	471	16	Q92NM0	Q92nm0 rhizobium m
12	39	72.2	477	5	P91348	P91348 caenorhabdi
13	38	70.4	174	16	Q92JL8	Q92jl8 rickettsia
14	38	70.4	281	5	Q9V3A9	Q9v3a9 drosophila
15	38	70.4	1186	3	Q12466	Q12466 saccharomyc
16	37	68.5	323	5	Q9GUN9	Q9gun9 caenorhabdi

```
070292
ID 070292
                PRELIMINARY;
                                PRT; 590 AA.
    070292;
AC
DT
    01-AUG-1998 (TrEMBLrel. 07, Created)
DT
    01-AUG-1998 (TrEMBLrel. 07, Last sequence update)
DT
    01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DΕ
    G PROTEIN-COUPLED RECEPTOR KINASE 5.
    GPRK5 OR GRK5.
GN
OS
    Mus musculus (Mouse)
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC 
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI_TaxID=10090;
```

RESULT

```
RN
     [1]
     SEQUENCE FROM N.A.
ΡP
     MEDLINE=99436149; PubMed=10506199;
P.X
     Premont R.T., Macrae A D , Aparicio S.A., Kendall H.E., Welch J.E.,
F:A
P.A
     Lefkowitz R J ,
     "The GRK4 subfamily of G protein-coupled receptor kinases. Alternative
PТ
F:T
     splicing, gene organization, and sequence conservation.";
     J. Biol. Chem. 274 29381-29389(1999).
-!- SIMILARITY BELONGS TO THE SER/THR FAMILY OF PROTEIN KINASES. GPRK
P.L
ac
CC
         SUBFAMILY
cc
     -!- SIMILARITY CONTAINS 1 RGS DOMAIN.
     EMBL; AF040746. AAC09267 1; -.
DR.
DP.
     HSSP; Q63450; 1A06
     MGD; MGI 109161; Gprk5
DΡ
DP.
     InterPro, IPR000719; Euk_pkinase.
     InterPro, IPR000239, GPCR_kinase.
DF.
DF
     InterPro: IPR000961, Pkinase_C.
     InterPro; IPR000342; RGS.
DF:
     InterPro: IPR002290, Ser thr pkinase.
DP.
DP.
     Pfam; PF00069; pkinase; 1.
     Pfam; PF00615; RGS: 1
DF:
     PRINTS; PR00717; GPCRKINASE.
DR
DP.
     SMART; SM00315, RGS; 1
     SMART; SM00220, S_TKc, 1
SMART; SM00133, S_TK_X, 1.
DR.
DR
     PROSITE, PS00107; PROTEIN_KINASE_ATP; 1.
     PROSITE, PS50011; PROTEIN_KINASE_DOM; 1.
DR
DR
     PROSITE; PS50132; RGS; 1
    ATP-binding, Kinase, Receptor; Transferase.
    SEQUENCE 590 AA; 67732 MW; F47D87397B1A2399 CRC64;
  Query Match 74.1%; Score 40; DB 11; Length 590; Best Local Similarity 85.7%; Pred. No. 40; Matches 6; Conservative 1; Mismatches 0; Indels
                                                     0; Indels 0; Gaps
        2 DDYGHLR 8
QУ
           11 1:
ďū
      320 DDYGHIR 326
PESULT
070297
ID
    070297
                  PRELIMINARY;
                                    PRT; 590 AA.
АĈ
     070297:
     01-AUG-1998 (TrEMBLrel. 07, Created)
DT
DT
     01-AUG-1998 (TrEMBLrel. 07, Last sequence update)
200
     01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE.
     G PROTEIN-COUPLED RECEPTOR KINASE 5.
     GPRK5 OR GRK5
GN
     Mus musculus (Mouse).
0S
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
00
     Mammalia, Eutheria, Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OC
OX
     NCBI TaxID=10090;
F.N
     [1]
P.P
     SEQUENCE FROM N A.
P.C
     STRAIN=129SVJ,
     MEDLINE=99436149; PubMed=10506199;
RX
PA
     Premont R.T., Macrae A.D., Aparicio S.A., Kendall H.E., Welch J.E.,
RА
     Lefkowitz R.J
RT
     "The GRE4 subfamily of G protein-coupled receptor kinases. Alternative
RT
     splicing, gene organization, and sequence conservation.";
\mathbf{EL}
     J. Biol. Chem 274 29381-29389(1999).
     -:- SIMILARITY BELONGS TO THE SER/THR FAMILY OF PROTEIN KINASES. GPRK
CC
CC
         SUBFAMILY
CC
      -!- SIMILARITY: CONTAINS 1 RGS DOMAIN.
     EMBL; AF040759; AAC09271 1; -.
DF:
DR.
     EMBL; AF040755; AAC09271 1; JOINED.
     EMBL; AF040756; AAC09271.1; JOINED.
DR.
     EMBL; AF040757; AAC09271 1; JOINED.
DF.
DR.
     EMBL; AF040758; AAC09271 1; JOINED.
```

```
DR HSSP; Q63450; 1A06.
DR.
      MGD; MGI:109161; Gprk5.
      InterPro; IPR000719; Euk_pkinase.
DR.
FR InterPro; IPR000239; GPCR_kinase.
\Gamma R
       InterPro; IPR000961; Pkinase_C.
      InterPro; IPR000342; RGS.
I:R
      InterPro; IPR002290; Ser_thr_pkinase.
\mathbb{D}\mathbb{R}
ER.
       Pfam; PF00069; pkinase; \overline{1}.
      Pfam; PF00615; RGS; 1.
DR.
DR.
      PRINTS; PR00717; GPCRKINASE.
      SMART; SM00315; RGS; 1.
SMART; SM00220; S_TKc; 1.
DR
DR
      SMART; SM00133; S_TK_X; 1.
PROSITE; PS00107; PROTEIN_KINASE_ATP; 1.
PROSITE; PS50011; PROTEIN_KINASE_DOM; 1.
DR
DR
DR PROSITE; PS50132; RGS; 1.
KW ATP-binding; Kinase; Receptor; Transferase.
SQ SEQUENCE 590 AA; 67796 MW; 22253281964DEF64 CRC64;
  Query Match 74.1%; Score 40; DB 11; Length 590; Best Local Similarity 85.7%; Pred. No. 40; Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps
          2 DDYGHLR 8
Qу
             320 DDYGHIR 326
```

9,9693746

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID: SSSPTA1649JXM

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
* * * * * *
                     Welcome to STN International
NEWS 1
                 Web Page URLs for STN Seminar Schedule - N. America
                 BLAST(R) searching in REGISTRY available in STN on the Web
NEWS 2 Jan 25
NEWS 3 Jan 29 FSTA has been reloaded and moves to weekly updates
NEWS 4 Feb 01 DKILIT now produced by FIZ Karlsruhe and has a new update
                 frequency
NEWS 5 Feb 19 Access via Tymnet and SprintNet Eliminated Effective 3/31/02
NEWS 6 Mar 08 Gene Names now available in BIOSIS
NEWS 7 Mar 22 TOXLIT no longer available
NEWS 8 Mar 22 TRCTHERMO no longer available
NEWS 9 Mar 28 US Provisional Priorities searched with P in CA/CAplus
                 and USPATFULL
NEWS 10 Mar 28 LIPINSKI/CALC added for property searching in REGISTRY
                PAPERCHEM no longer available on STN. Use PAPERCHEM2
NEWS 11 Apr 02
instead.
NEWS 12 Apr 08
                "Ask CAS" for self-help around the clock
                 BEILSTEIN: Reload and Implementation of a New Subject Area
NEWS 13 Apr 09
                 ZDB will be removed from STN
NEWS 14 Apr 09
NEWS 15 Apr 19 US Patent Applications available in IFICDB, IFIPAT, and
IFIUDB
NEWS 16 Apr 22 Records from IP.com available in CAPLUS, HCAPLUS, and
ZCAPLUS
NEWS 17 Apr 22
                 BIOSIS Gene Names now available in TOXCENTER
NEWS 18 Apr 22
                 Federal Research in Progress (FEDRIP) now available
NEWS 19
        Jun 03
                New e-mail delivery for search results now available
NEWS 20 Jun 10 MEDLINE Reload
NEWS 21 Jun 10
                PCTFULL has been reloaded
NEWS 22
        Jul 02
                FOREGE no longer contains STANDARDS file segment
NEWS 23 Jul 19 NTIS to be reloaded July 28, 2002
NEWS 24 Jul 22 USAN to be reloaded July 28, 2002;
                 saved answer sets no longer valid
NEWS EXPRESS February 1 CURRENT WINDOWS VERSION IS V6.0d,
              CURRENT MACINTOSH VERSION IS V6.0a(ENG) AND V6.0Ja(JP),
              AND CURRENT DISCOVER FILE IS DATED 05 FEBRUARY 2002
NEWS HOURS
              STN Operating Hours Plus Help Desk Availability
NEWS INTER
              General Internet Information
NEWS LOGIN
              Welcome Banner and News Items
NEWS PHONE
              Direct Dial and Telecommunication Network Access to STN
              CAS World Wide Web Site (general information)
NEWS WWW
```

Enter NEWS followed by the item number or name to see news on that specific topic.

All use of STN is subject to the provisions of the STN Customer agreement. Please note that this agreement limits use to scientific

research. Use for software development or design or implementation of commercial greways or other similar uses is bhibited and may result in loss of user privileges and other penalties.

FILE 'HOME' ENTERED AT 10:00:34 ON 24 JUL 2002

=> file medline biosis embase caplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL

ENTRY SESSION

FULL ESTIMATED COST

0.21

0.21

FILE 'MEDLINE' ENTERED AT 10:00:47 ON 24 JUL 2002

FILE 'BIOSIS' ENTERED AT 10:00:47 ON 24 JUL 2002 COPYRIGHT (C) 2002 BIOLOGICAL ABSTRACTS INC.(R)

FILE 'EMBASE' ENTERED AT 10:00:47 ON 24 JUL 2002

COPYRIGHT (C) 2002 Elsevier Science B.V. All rights reserved.

FILE 'CAPLUS' ENTERED AT 10:00:47 ON 24 JUL 2002

USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.

PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

=> s dmgpcr

1.1 1 DMGPCR

=> s drosophila (p) receptor (p) coupled (p) bind

51 DROSOPHILA (P) RECEPTOR (P) COUPLED (P) BIND 1.2

=> dup rem 12

PROCESSING COMPLETED FOR L2

18 DUP REM L2 (33 DUPLICATES REMOVED)

=> d l3 total ibib kwic

ANSWER 1 OF 18 MEDLINE DUPLICATE 1

ACCESSION NUMBER: 2002045994

IN-PROCESS

DOCUMENT NUMBER:

21630175 PubMed ID: 11754840

TITLE:

Regulation of the Rhodopsin Protein Phosphatase, RDGC,

through Interaction with Calmodulin.

AUTHOR:

SOURCE:

Lee S J; Montell C

CORPORATE SOURCE: Department of Biological Chemistry and, Department of Neuroscience, The Johns Hopkins University School of

Medicine, 21205, Baltimore, MD, USA.

NEURON, (2001 Dec 20) 32 (6) 1097-106.

Journal code: 8809320. ISSN: 0896-6273.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE:

Entered STN: 20020124

Last Updated on STN: 20020124

Hundreds of G protein-coupled receptors (GPCRs) and at

least six GPCR kinases have been identified, but the only GPCR phosphatase

that has been definitively demonstrated is the rhodopsin phosphatase encoded by to rdgC locus of **Drosophila**. Muta ons in rdgC result in defects in termination of the light response and cause severe retinal degeneration. In the current work, we demonstrate that RDGC binds to calmodulin, and a mutation in an IQ motif that eliminates the calmodulin/RDGC interaction prevents dephosphorylation of rhodopsin

in

vivo. . .

L3 ANSWER 2 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 2001:244124 BIOSIS DOCUMENT NUMBER: PREV200100244124

TITLE: The 1.8ANG crystal structure of InaD PDZ1 complexed with

its peptide target reveals a novel mode of PDZ domain

binding.

AUTHOR(S): Pliske, Michelle (1); Sondek, John (1)

CORPORATE SOURCE: (1) Biochemistry and Biophysics, UNC-Chapel Hill, Mary

Ellen Jones Bldg., Chapel Hill, NC, 27599 USA

SOURCE: FASEB Journal, (March 8, 2001) Vol. 15, No. 5, pp. A723.

print.

Meeting Info.: Annual Meeting of the Federation of

American

Societies for Experimental Biology on Experimental Biology

2001 Orlando, Florida, USA March 31-April 04, 2001

ISSN: 0892-6638.

DOCUMENT TYPE: Conference LANGUAGE: English SUMMARY LANGUAGE: English

AB **Drosophila** phototransduction is a model system for the study of G-protein **coupled** phospholipase-C (PLC) signaling pathways in

complex organisms. In this cascade light activates the

seven-transmembrane

receptor rhodopsin, which in turn activates Gq, allowing its dissociation into signaling-competent alpha and betagamma subunits. Gqalpha induces the PLC-beta4 homolog no receptor potential A (norpA) to cleave phosphatidylinositol-4,5-bisphosphate (PIP2) to the second messengers inositol tri-phosphate (IP3) and diacylglycerol (DAG), leading to the. . . multi-domain scaffolding protein inactivation no after-potential D (inaD). InaD contains five tandem PDZ protein interaction domains, each of which can bind one or multiple phototransduction proteins. PDZ domains bind to the extreme carboxy-terminal (C-terminal) three amino acids of their targets, including the free carboxyl group. InaD PDZ1 binds to norpA, whose C-terminal sequence is FCA. The X-ray crystal structure of PDZ1 bound to a norpA C-terminal heptapeptide was. . .

L3 ANSWER 3 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 2001:492061 BIOSIS DOCUMENT NUMBER: PREV200100492061

TITLE: Comparison of adenylyl cyclase stimulation by 5-HT4(b) and

5-HT7(a) receptors using the Ecdysone-Inducible Mammalian

Expression System.

AUTHOR(S): Bruheim, S. (1); Andressen, K. W. (1); Krobert, K. A. (1);

Levy, F. O. (1)

CORPORATE SOURCE: (1) MSD Cardiovascular Res. Ctr. and Dept. of Pharmacol.,

Univ. of Oslo, Oslo Norway

SOURCE: Society for Neuroscience Abstracts, (2001) Vol. 27, No. 1,

pp. 690. print.

Meeting Info.: 31st Annual Meeting of the Society for Neuroscience San Diego, California, USA November 10-15,

2001

ISSN: 0190-5295.

DOCUMENT TYPE: Conference LANGUAGE: English SUMMARY LANGUAGE: English

AB The serotonin (5-HT) receptors 5-HT4 and 5-HT7 are G-protein

coupled receptors that activate adenylyl cyclase (AC) and exist in everal splice variants differing only in their intracellular

carboxyl terminal tails. We wanted to determine if activation of AC differed between the 5-HT4(b) and 5-HT7(a) receptors. Comparison of receptor function using constitutive expression systems can be confounded by different receptor expression levels and clonal cell line differences. By using the Ecdysone-Inducible Mammalian Expression System we could reproducibly express varying levels of receptor in the same clonal cell line. This system utilizes a heterodimer (VgRxR) of the modified ecdysone receptor (VgEcR) from Drosophila and the retionoid X receptor (RXR). This receptor binds a hybrid ecdysone response element (E/GRE) in the presence of the synthetic analog of ecdysone, ponasterone A. HEK293 cells stably expressing the heterodimer VgRxR receptor were stably transfected with a vector containing the coding regions for 5-HT4(b) and 5-HT7(a) receptors downstream of the E/GRE. Radioligand binding revealed low constitutive expression of both receptors, which could be titrated up to 3.7 pmol/mg protein with ponasterone A. Preliminary data indicate that constitutive AC activity

and

potency (EC50) of 5-HT are **receptor** level dependent at the 5-HT4(b) **receptor** but not at the 5-HT7(a) **receptor**. Additionally, the 5-HT7(a) **receptor** activated AC more efficiently than the 5-HT4(b) **receptor** over a wide range of expression levels. Comparative studies on inverse agonism are ongoing.

L3 ANSWER 4 OF 18 MEDLINE DUPLICATE 2

ACCESSION NUMBER: 2001271685

0001271685 MEDLINE

DOCUMENT NUMBER:

21261850 PubMed ID: 11369206

TITLE:

Mutations in the sterol-sensing domain of Patched suggest

role for vesicular trafficking in Smoothened regulation.

COMMENT:

Erratum in: Curr Biol 2001 Jul 24;11(14):1153

AUTHOR:

Strutt H; Thomas C; Nakano Y; Stark D; Neave B; Taylor A

Μ;

а

Ingham P W

CORPORATE SOURCE:

MRC Intercellular Signalling Group, Centre for

Developmental Genetics, Department of Biomedical Science,

University of Sheffield, Sheffield, United Kingdom.

SOURCE:

CURRENT BIOLOGY, (2001 Apr 17) 11 (8) 608-13.

Journal code: 9107782. ISSN: 0960-9822.

PUB. COUNTRY:

England: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200108

ENTRY DATE:

Entered STN: 20010813

Last Updated on STN: 20010813 Entered Medline: 20010809

AB The tumor suppressor gene patched (ptc) encodes an approximately 140 kDa polytopic transmembrane protein [1-3] [corrected] that **binds** members of the Hedgehog (Hh) family of signaling proteins [4-6] [corrected] and regulates the activity of Smoothened (Smo), a G protein-coupled receptor-like protein essential for Hh signal

coupled receptor-like protein essential for Hh signal transduction [7-9] [corrected]. Ptc contains a sterol-sensing domain

(SSD)

[10, 11] [corrected], a motif found. . . (Hh) signaling by facilitating

the regulated secretion and sequestration of the Hh protein [16] [corrected], to which it is covalently **coupled**. In addition, cholesterol synthesis inhibitors block the ability of cells to respond to Hh [18, 19] [corrected], and this finding. . . has so far been

lacking.

Here we describe the identification and characterization of two missense mutations in the SSD of **Drosophila** Ptc; strikingly, while both

mutations abolish Smo repression, neither affects the ability of Ptc to interact with the We speculate. . .

L3 ANSWER 5 OF 18 MEDLINE DUPLICATE 3

ACCESSION NUMBER: 2002053834

002053834 MEDLINE

DOCUMENT NUMBER:

21638057 PubMed ID: 11779634

TITLE:

The transcription factors Sp1 and Sp3 are required for human angiotensin II type 1 receptor gene expression in

H295-R cells.

AUTHOR:

Zhao X; Martin M M; Elton T S

CORPORATE SOURCE:

Department of Chemistry and Biochemistry, Brigham Young University, C206 Benson Building, P.O. Box 25700, Provo,

UT

84602-5700, USA.

CONTRACT NUMBER:

HL48848 (NHLBI)

SOURCE:

BIOCHIMICA ET BIOPHYSICA ACTA, (2001 Dec 30) 1522 (3)

195-206.

Journal code: 0217513. ISSN: 0006-3002.

PUB. COUNTRY:

Netherlands

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200202

ENTRY DATE:

Entered STN: 20020125

Last Updated on STN: 20020222 Entered Medline: 20020221

AB . . . peptide hormone angiotensin II regulates a variety of

physiological responses which are mediated by its interaction with high affinity G protein-coupled receptors localized on the

surface of target cells. Our previous studies have demonstrated that a

145

bp sequence within the promoter region was required for basal level expression of the human angiotensin II type 1 receptor (hAT(1)R) gene. In the present study, deletional analysis of the hAT(1)R promoter localized the major regulatory sequence to two overlapping. . .

binding

site for Sp1 prevented the formation of the DNA-protein complexes. Supershift EMSAs also demonstrated that Sp1 and Sp3 could **bind** to the GC boxes present within the -105 to -85 bp region of the hAT(1)R promoter. Transactivation experiments utilizing **Drosophila** SL2 cells, which lack endogenous Sp family transcription factors,

demonstrated

that Sp1 and Sp3 activated the hAT(1)R promoter and that. . .

L3 ANSWER 6 OF 18 MEDLINE

DLINE DUPLICATE 4

ACCESSION NUMBER:

CORPORATE SOURCE:

2001688270

MEDLINE

DOCUMENT NUMBER:

21592298 PubMed ID: 11734218

TITLE:

Identification of mouse trp homologs and lipid rafts from

spermatogenic cells and sperm.

AUTHOR:

Trevino C L; Serrano C J; Beltran C; Felix R; Darszon A Department of Genetics and Molecular Physiology, Institute

of Biotechnology, UNAM, Cuernavaca, Mexico. FEBS LETTERS, (2001 Nov 30) 509 (1) 119-25.

SOURCE:

Journal code: 0155157. ISSN: 0014-5793.

their protein products (Trp1, Trp3 and Trp6) in mouse sperm. Allegedly

PUB. COUNTRY:

Netherlands

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT: Priority Journals

ENTRY MONTH:

200201

ENTRY DATE:

Entered STN: 20011206

Last Updated on STN: 20020125 Entered Medline: 20020122

AB

. . . of the membrane systems that regulate Ca(2+) in sperm. In this report, we provide evidence for the expression of seven **Drosophila** transient **receptor** potential homolog genes (trp1-7) and three of

some trps. . . . major component of caveolae, a subset of lipid rafts potentially cortant for signaling events and Ta(2+) flux. Furthermore, by using fluorescein-coupled cholera toxin B subunit, which specifically binds to the raft component ganglioside GM1, we identified caveolin- and Trp-independent lipid rafts residing in the plasma membrane of mature. . .

L3 ANSWER 7 OF 18 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1999:297449 CAPLUS

DOCUMENT NUMBER: 130:321591

TITLE: Cloning and cDNA sequence of an invertebrate

octopamine receptor

INVENTOR(S): Davis, Ronald L.; Han, Kyung-an; Millar, Neil S.

PATENT ASSIGNEE(S): Baylor College of Medicine, USA

SOURCE: PCT Int. Appl., 58 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE WO 9921891 A1 19990506 WO 1998-US22808 19981027 -----W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG AU 1999-19008 19981027 AU 9919008 A1 19990517 US 1997-63391P P 19971027 PRIORITY APPLN. INFO.: WO 1998-US22808 W 19981027 REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

The present invention provides a novel octopamine receptor comprised of an invertebrate receptor which binds octopamine and is coupled to adenylyl cyclase system. The invention also includes methods of using the octopamine receptor to screen for agonists, antagonists and pesticides. In this method the octopamine receptor is inserted into invertebrate or vertebrate cells, test compd. is added and the activity of the octopamine receptor coupled to the adenylyl cyclase system or the internal Ca2+ system is measured. Also included is an expression system for prodn. of the octopamine receptor. The octopamine receptor cDNA was cloned from Drosophila melanogaster using PCR and single-strand conformation polymorphism.

L3 ANSWER 8 OF 18 MEDLINE DUPLICATE 5

ACCESSION NUMBER: 1999147062 MEDLINE

DOCUMENT NUMBER: 99147062 PubMed ID: 10022914

TITLE: Identification of a novel family of targets of PYK2

related

to Drosophila retinal degeneration B (rdgB) protein.

AUTHOR: Lev S; Hernandez J; Martinez R; Chen A; Plowman G;

Schlessinger J

CORPORATE SOURCE: Sugen, Inc., South San Francisco, California 94080, USA. SOURCE: MOLECULAR AND CELLULAR BIOLOGY, (1999 Mar) 19 (3) 2278-88.

Journal code: 8109087. ISSN: 0270-7306.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: ENTRY DATE: 199903

Entered STN: 19990402

Last Updated on STN: 19990402

Entered Medline: 19990325

AB The protein tyrosine kinase PYK2 has been implicated in signaling pathways

activated by G-protein-coupled receptors,

intracellular calcium, and stress signals. Here we describe the molecular cloning and characterization of a novel family of PYK2-binding proteins designated Nirs (PYK2 N-terminal domain-interacting receptors).

The three Nir proteins (Nir1, Nir2, and Nir3) **bind** to the amino-terminal domain of PYK2 via a conserved sequence motif located in the carboxy terminus. The primary structures of. . . region homologous to phosphatidylinositol (PI) transfer protein, and an acidic domain. The Nir proteins are the human homologues of the **Drosophila** retinal degeneration B protein (rdgB), a protein implicated in the visual transduction pathway in flies. We demonstrate that Nirs are. . .

family

of evolutionarily conserved PYK2-binding proteins that play a role in the control of calcium and phosphoinositide metabolism downstream of G-protein-coupled receptors.

L3 ANSWER 9 OF 18 MEDLINE DUPLICATE 6

ACCESSION NUMBER: 1998227978 MEDLINE

DOCUMENT NUMBER: 98227978 PubMed ID: 9569023

TITLE: Disabled-2 (Dab2) is an SH3 domain-binding partner of

Grb2.

AUTHOR: Xu X X; Yi T; Tang B; Lambeth J D

CORPORATE SOURCE: Department of Biochemistry, and Winship Cancer Center,

Emory University School of Medicine, Atlanta, Georgia

30322, USA.

CONTRACT NUMBER: R 01 CA75389-01 (NCI)

R01CA46508 (NCI)

SOURCE: ONCOGENE, (1998 Mar 26) 16 (12) 1561-9.

Journal code: 8711562. ISSN: 0950-9232.

PUB. COUNTRY: ENGLAND: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199805

ENTRY DATE: Entered STN: 19980520

Last Updated on STN: 20000303 Entered Medline: 19980513

Disabled-2 (Dab2), a mammalian structural homolog of Drosophila AΒ Disabled (Dab), is a mitogen-responsive phosphoprotein. It has been speculated to be a negative regulator of growth since its expression. exchange factor for Ras. The proline-rich sequences of Sos mediate the interaction of Sos with Grb2, an adaptor protein which coupled tyrosine kinase receptors to Sos. Herein, we have investigated the possibility that Dab2 interacts with Grb2. In experiments of co-immunoprecipitation from BAC1.2F5 macrophage. . . disrupting the Grb2-Sos complex. The expressed proline-rich domain of Dab2 (#600-730) bound Grb2, but other regions of Dab2 failed to bind Grb2. Both of the individual SH3 domains of Grb2 bound to Sos (N-terminal SH3 domain >> C-terminal SH3 domain), but. . . to Dab2 required the intact Grb2, suggesting cooperative binding using both SH3 domains of Grb2. These data indicate that Dab2 binds to the SH3 domains of Grb2 via its C-terminal proline-rich sequences. Dab2 may modulate growth factor/Ras pathways by competing with.

3 ANSWER 10 OF 18 MEDLINE DUPLICATE 7

ACCESSION NUMBER: 1998340528 MEDLINE

DOCUMENT NUMBER: 98340528 PubMed ID: 9675877

TITLE: The c-Cbl oncoprotein.

AUTHOR: Lupher M L Jr; Andoniou C E; Bonita D; Miyake S; Band H CORPORATE SOURCE: Department of Medicine, Brigham and Women's Hospital,

Harvard Medical School, Boston, MA 02115, USA. INTERNATIONAL JOURNAL OF BIOCH STRY AND CELL SOURCE:

STRY AND CELL BIOLOGY,

(1998 Apr) 30 (4) 439-44. Ref: 16

Journal code: 9508482. ISSN: 1357-2725.

PUB. COUNTRY: ENGLAND: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE:

English

Priority Journals FILE SEGMENT:

ENTRY MONTH:

199808

Entered STN: 19980820 ENTRY DATE:

> Last Updated on STN: 19980820 Entered Medline: 19980813

Cbl has emerged as a novel signal transducing protein downstream of a AΒ number of cell surface receptors coupled to tyrosine

kinases. Identified as the protein product of the c-cbl proto-oncogene, the cellular homolog to the transforming gene of. . . finger render

Cbl

oncogenic, whereas wild type Cbl is non-transforming, even if overexpressed. Cbl serves as a substrate of both receptor and non-receptor tyrosine kinases, and binds to adaptor proteins Grb2, Crk and the p85 subunit of PI-3-kinase. Additionally, both Caenorhabditis elegans and Drosophila Cbl homologs, SLI-1 and D-Cbl, respectively, have been identified as negative regulators of the LET-23/DER receptor tyrosine kinases. Finally, oncogenic mutants of Cbl, when expressed in fibroblasts, upregulate the signaling cascade downstream of the platelet-derived growth factor receptor alpha in a Cbl-PTB domain-dependent manner. Together, these findings position Cbl as a central player in the regulation of tyrosine.

ANSWER 11 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 1999:340756 BIOSIS DOCUMENT NUMBER:

PREV199900340756

TITLE:

SOURCE:

First Annual Jorge Chevesich Memorial Lecture: A

supramolecular signaling complex required for Drosophila

visual transduction.

AUTHOR(S):

Montell, Craig (1)

CORPORATE SOURCE:

(1) Departments of Biological Chemistry and Neuroscience, Johns Hopkins University School of Medicine, 725 N. Wolfe

Street, Baltimore, MD, 21205 USA Einstein Quarterly Journal of Biology and Medicine, (1998)

Vol. 15, No. 4, pp. 198-211.

ISSN: 0724-6706.

DOCUMENT TYPE: General Review

LANGUAGE: English SUMMARY LANGUAGE: English

Drosophila phototransduction represents one of the fastest known G-protein coupled signaling cascades. Exposure of the photoreceptor cells to light leads to activation of the light-induced cation influx channels, TRP and. . . in phototransduction are linked into a supramolecular signaling complex (signalplex) has led to a reevaluation of the mechanisms underlying the Drosophila photoresponse. The central player is INAD, a protein with five protein interaction motifs referred to as PDZ domains. At least seven signaling molecules bind to INAD. These include rhodopsin, phospholipase C-beta, protein kinase C, TRP, TRPL, calmodulin and an unconventional myosin, NINAC. Someof the. . . Since more than five proteins interact with INAD, it would appear that a single INAD monomer lacks the capacity to bind to each of its targets simultaneously. The finding that INAD is capable of forming homo-multimers in vitro raises the possibility that the entire phototransduction cascade may be physically coupled in the signalplex. Nearly all of the proteins that function in the signalplex have known vertebrate homologs. These include a. . neurons of the central nervous system around the time of birth. TRPC3 appears to be activated through stimulation of the receptor

tyrosine kinase, TrkB, and phospholipase C-gamma. Thus, a variety of g to the stimulation of phosph pathways lea pase C appear to.

ANSWER 12 OF 18 MEDLINE DUPLICATE 8

ACCESSION NUMBER: 1998024192

MEDLINE

DOCUMENT NUMBER:

98024192 PubMed ID: 9356510

TITLE:

Association of INAD with NORPA is essential for controlled

activation and deactivation of Drosophila

phototransduction

CORPORATE SOURCE:

in vivo.

AUTHOR:

Shieh B H; Zhu M Y; Lee J K; Kelly I M; Bahiraei F Department of Pharmacology, Vanderbilt University,

Nashville, TN 37232-6600, USA.. shiehb@ctrvax.vanderbilt.edu

CONTRACT NUMBER:

EY09743 (NEI)

SOURCE:

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1997 Nov 11) 94 (23) 12682-7.

Journal code: 7505876. ISSN: 0027-8424.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199712

ENTRY DATE:

Entered STN: 19980109

Last Updated on STN: 19980109 Entered Medline: 19971216

Visual transduction in Drosophila is a G protein-coupled phospholipase C-mediated process that leads to depolarization via activation of the transient receptor potential (TRP) calcium channel. Inactivation-no-afterpotential D (INAD) is an adaptor protein containing PDZ domains known to interact with TRP. Immunoprecipitation studies indicate that INAD also binds to eye-specific protein kinase C and the phospholipase C, no-receptor-potential A (NORPA). By overlay assay and site-directed mutagenesis we have defined the essential elements of the NORPA-INAD association and identified.

ANSWER 13 OF 18 MEDLINE 1.3

DUPLICATE 9

ACCESSION NUMBER: 97197784

MEDLINE

DOCUMENT NUMBER:

PubMed ID: 9045634 97197784

TITLE:

Cloning and expression of a complementary DNA encoding a molluscan octopamine receptor that couples to chloride

channels in HEK293 cells.

AUTHOR:

Gerhardt C C; Lodder H C; Vincent M; Bakker R A; Planta R

J; Vreugdenhil E; Kits K S; van Heerikhuizen H

CORPORATE SOURCE:

Department of Biochemistry and Molecular Biology, Research Institute Neurosciences, Vrije Universiteit, De Boelelaan

1083, 1081 HV Amsterdam, The Netherlands.

SOURCE:

JOURNAL OF BIOLOGICAL CHEMISTRY, (1997 Mar 7) 272 (10)

6201-7.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT: OTHER SOURCE: Priority Journals GENBANK-U62770

ENTRY MONTH:

199704

ENTRY DATE:

Entered STN: 19970424

Last Updated on STN: 20000303 Entered Medline: 19970414

A cDNA encoding a G-protein-coupled receptor was AΒ

cloned from the central nervous system of the pond snail Lymnaea stagnalis. The predicted amino acid sequence of this cDNA most closely resembles the Drosophila tyramine/octopamine receptor,

the Locusta tyramine receptor, and an octopamine

receptor (Lym oal) that we recently cloned from Lymnaea. After

stable expression of the cDNA in HEK293 cells, we found that [3H]rauwolsd binds with high affinity to treceptor (KD = 6.2.10(-9) M). Octopamine appears to be the most potent naturally occurring agonist to displace the [3H]rauwolscine binding (Ki = 3.0.10(-7)

M). Therefore, the **receptor** is considered to be an octopamine **receptor** and is consequently designated Lym oa2. The novel **receptor** shares little pharmacological resemblance with Lym oa1, indicating that the two **receptors** represent different octopamine **receptor** subfamilies. Octopaminergic stimulation of Lym oa2 does not induce changes in intracellular concentrations of cAMP or inositol phosphates. However, electrophysiological. . .

DUPLICATE 10

L3 ANSWER 14 OF 18 MEDLINE

ACCESSION NUMBER: 97472416 MEDLINE

DOCUMENT NUMBER: 97472416 PubMed ID: 9333241

TITLE: Prolonged photoresponses in transgenic mouse rods lacking

arrestin.

AUTHOR: Xu J; Dodd R L; Makino C L; Simon M I; Baylor D A; Chen J

CORPORATE SOURCE: Division of Biology, California Institute of Technology,

Pasadena 91125, USA.

SOURCE: NATURE, (1997 Oct 2) 389 (6650) 505-9.

Journal code: 0410462. ISSN: 0028-0836.

PUB. COUNTRY: ENGLAND: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199710

ENTRY DATE: Entered STN: 19971105

Last Updated on STN: 19971105 Entered Medline: 19971022

AB Arrestins are soluble cytoplasmic proteins that **bind** to G-protein-**coupled receptors**, thus switching off activation of the G protein and terminating the signalling pathway that triggers the cellular response. Although visual. . . was halved, indicating that arrestin binding is not rate limiting for recovery of the rod photoresponse, as it is in **Drosophila**. With arrestin absent, the flash response displayed a rapid partial recovery followed by a prolonged final phase. This behaviour indicates. . .

L3 ANSWER 15 OF 18 MEDLINE DUPLICATE 11

ACCESSION NUMBER: 97347296 MEDLINE

DOCUMENT NUMBER: 97347296 PubMed ID: 9203635

TITLE: Molecular cloning and pharmacological characterization of

а

molluscan octopamine receptor.

AUTHOR: Gerhardt C C; Bakker R A; Piek G J; Planta R J;

Vreugdenhil

E; Leysen J E; Van Heerikhuizen H

CORPORATE SOURCE: Department of Biochemistry and Molecular Biology, Research

Institute Neurosciences, Vrije Universiteit, Amsterdam,

The

Netherlands.

SOURCE: MOLECULAR PHARMACOLOGY, (1997 Feb) 51 (2) 293-300.

Journal code: 0035623. ISSN: 0026-895X.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199707

ENTRY DATE: Entered STN: 19970805

Last Updated on STN: 19970805 Entered Medline: 19970723

AB We describe the cloning and functional expression of a cDNA encoding a novel G protein-coupled receptor, which was isolated from the central nervous system of the pond snail Lymnaea stagnalis. The

amino acid sequence predicted by this cDNA shows highest similarity with the sequence the Locusta tyramine receptor he Drosophila tyramine/octopamine receptor, and the mammalian alpha-adrenergic receptors. On expression in mammalian cells, [3H]rauwolscine, an alpha2-adrenergic receptor antagonist, binds with high affinity  $(K(D) = 2.9 \times 10(-9) M)$  to the receptor. Of several tested neurotransmitters, octopamine (which is considered to be the invertebrate counterpart of norepinephrine) showed the highest affinity  $(1.9 \times 10(-6) \text{ M})$  for the **receptor**. Therefore, we consider this receptor to be the first true octopamine receptor to be cloned. The ligand binding properties of the novel receptor, designated Lym oal, seem to be distinct from any of the binding profiles described for octopamine receptors in tissue preparations. Although the pharmacological profile of Lym oal shows some similarity with that of Tyr/Oct-Dro and Tyr-Loc, there. . . also clear differences. In particular, phentolamine, chlorpromazine, and mianserine display markedly higher affinities for Lym oal than for the insect receptors. As far as the vertebrate adrenergic receptors are concerned, the ligand binding properties of Lym oal resemble alpha2-adrenergic receptors more than they do alpha- or beta-adrenergic receptors. Octopaminergic stimulation of Lym oal induces an increase in both inositol phosphates and cAMP (EC50 =  $9.1 \times 10(-7) \text{ M}$ . . .  $\times 10(-6) \text{ M}$ , respectively). This is in contrast to the signal transduction pathways described for the related tyramine- and alpha2-adrenergic receptors, which couple in an inhibitory way to adenylyl cyclase. ANSWER 16 OF 18 MEDLINE DUPLICATE 12 ACCESSION NUMBER: 91198639 MEDLINE PubMed ID: 1849770 DOCUMENT NUMBER: 91198639 TITLE: Very high affinity interaction of DPI 201-106 and BDF 8784 enantiomers with the phenylalkylamine-sensitive Ca2(+)-channel in Drosophila head membranes. Glossmann H; Zech C; Striessnig J; Staudinger R; Hall L; AUTHOR: Greenberg R; Armah B I CORPORATE SOURCE: Institut fur Biochemische Pharmakologie, Innsbruck, Austria. SOURCE: BRITISH JOURNAL OF PHARMACOLOGY, (1991 Feb) 102 (2) 446-52. Journal code: 7502536. ISSN: 0007-1188. PUB. COUNTRY: ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English FILE SEGMENT: Priority Journals ENTRY MONTH: 199105 ENTRY DATE: Entered STN: 19910607 Last Updated on STN: 19970203 Entered Medline: 19910523 1. Piperazinylindoles (DPI 201-106, BDF 8784), drugs known to act on AB voltage-dependent Na(+)-channels, bind with very high affinity to a Ca2(+)-channel-associated phenylalkylamine receptor in Drosophila melanogaster head membranes. These compounds and (+)-tetrandrine, a naturally occurring Ca2(+)-antagonist, were the most selective inhibitors for phenylalkylamine-labelled Drosophila Ca2(+)-channels compared to mammalian L-type Ca2(+)-channels. 2.

Replacement of the cyano group by a methyl group in (+)-DPI 201-106 ((+)-BDF 8784) increases the IC50 value for inhibition of phenylalkylamine

labelling of Drosophila Ca2(+)-channels from 0.29 to 2.1 nM but decreases the IC50 value for inhibition of phenylalkylamine labelling of mammalian skeletal muscle. . . to 49 nM. 3. DPI 201-106 enantiomers completely block (at 0.1 microM) phenylalkylamine photolabelling of a 136 K polypeptide in **Drosophila** head membranes whereas 10 microM aconitine or lidocaine are without effect. 4. Assessment of the

Ca2(+)-antagonist effects of the substituted. . . and chemical selectivity elated to local anaesthetic addity. 6. It is proposed that a very high affinity piperazinylindole-selective site is coupled to the phenylalkylamine receptor of Drosophila Ca2(+)-channels. These sites are still present on mammalian L-type Ca2(+)-channels but have lower affinity and/or are less tightly coupled to phenylalkylamine receptors on the alpha 1-subunit.

ANSWER 17 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 1989:224365 BIOSIS

BA87:115982 DOCUMENT NUMBER:

TITLE:

STRUCTURE-MEMBRANE MECHANISM OF DISTANT REGULATION OF

INTERCELLULAR GAP JUNCTION PERMEABILITY.

AUTHOR(S):

MAZHUL' V M; KONEV S V; YANCHEVSKAYA T G; FININ V S CORPORATE SOURCE: INST. PHOTOBIOL., ACAD. SCI. B. SSR, MINSK, USSR. SOURCE: BIOFIZIKA, (1989) 33 (6), 1023-1028.

CODEN: BIOFAI. ISSN: 0006-3029.

FILE SEGMENT:

BA; OLD Russian

LANGUAGE:

. . shown by ESR and tryptophane fluorescence at room temperature that concanavalin A (Con A) at a concentration of 10 mq/ml binds with glycoprotein receptors on the surface of salivary gland cells of Drosophila virilis larva beyong gap junction regions initiating generalized structural reorganization of plasmic membranes. The reorganization is coupled with a decrease in permeability of intercellular channels to small inorganic ions and molecules of the organic dye fluorescein. Treatment. . . structural and functional effects were reversible and nullified by substituting 4 mM .alpha.-D-glucopyronoside for lecitin in cell surface receptors. The obtained results suggest the existence of a structural membrane mechanism of distant regulation of intercellular communications according to the following pattern:.

ANSWER 18 OF 18 CAPLUS COPYRIGHT 2002 ACS L3

ACCESSION NUMBER: 1989:92520 CAPLUS

DOCUMENT NUMBER:

110:92520

TITLE:

Structure-membrane mechanism of distant regulation of

intercellular gap junction permeability

AUTHOR(S):

Mazhul, V. M.; Konev, S. V.; Yanchevskaya, T. G.;

Phinin, V. S.

CORPORATE SOURCE:

SOURCE:

Inst. Photobiol., Minsk, USSR Biofizika (1988), 33(6), 1023-8 CODEN: BIOFAI; ISSN: 0006-3029

DOCUMENT TYPE:

Journal Russian

LANGUAGE:

It was shown by ESR and tryptophan phosphorescence at room temp. that  ${\tt Con}$ AB A at 10 mg/mL binds with glycoprotein receptors on the surface of salivary gland cells of Drosophila virilis larva beyond gap junction regions initiating generalized structural reorganization of plasma membranes. The reorganization is coupled with a decrease in permeability of intercellular channels to small inorg. ions and fluorescein. Treatment of cells with dimethylsuberimidate-HCl,

reagent producing intra- and interprotein links which stabilize the network of plasma membranes, blocked the capacity of Con A to initiate structural reorganization of the membranes and disrupt intercellular communications. Con A-induced structural and functional effects were abolished by addn. of 4 mM .alpha.-D-glucopyranoside, which displaced the lectin on the cell surface receptors. The obtained results suggest the existence of a structural membrane mechanism of distal regulation of intercellular communications according to the following pattern: local structural reorganizations initiated beyond gap junction regions, generalization of the structural reorganization over the protein network of plasma membranes, involvement of high-permeability contact membranes in the reorganization, change in the structural organization

а

joining of protein half-channels of gap junctions, and modification of intercellula hannel permeability of small in g. ions and low-mol.-wt. org. compds.

=> log y

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	25.83	26.04
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL SESSION
CA SUBSCRIBER PRICE	-1.24	-1.24

STN INTERNATIONAL LOGOFF AT 10:02:08 ON 24 JUL 2002